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Genome sequencing and analysis of the filamentous fungus *Penicillium chrysogenum*

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Industrial penicillin production with the filamentous fungus *Penicillium chrysogenum* is based on an unprecedented effort in microbial strain improvement. To gain more insight into penicillin synthesis, we sequenced the 32.19 Mb genome of *P. chrysogenum* Wisconsin54-1255 and identified numerous genes responsible for key steps in penicillin production. DNA microarrays were used to compare the transcriptomes of the sequenced strain and a penicillinG high-producing strain, grown in the presence and absence of the side-chain precursor phenylacetic acid. Transcription of genes involved in biosynthesis of valine, cysteine and α -aminoadipic acid—precursors for penicillin biosynthesis—as well as of genes encoding microbody proteins, was increased in the high-producing strain. Some gene products were shown to be directly controlling β -lactam output. Many key cellular transport processes involving penicillins and intermediates remain to be characterized at the molecular level. Genes predicted to encode transporters were strongly overrepresented among the genes transcriptionally upregulated under conditions that stimulate penicillinG production, illustrating potential for future genomics-driven metabolic engineering.

Penicillins and derived β -lactam antibiotics have dramatically transformed health care and quality of life in the 80 years since Fleming's discovery of *Penicillium* that produces penicillins¹. Large-scale production of β -lactam antibiotics is the result of sustained industrial strain improvement, representing numerous rounds of mutagenesis and selection. Although information on industrial processes is proprietary, product titers and productivities have increased by at least three orders of magnitude in the past 60 years², representing an unprecedented success in classical strain improvement.

Current industrial strains are derived from a single natural isolate of *P. chrysogenum*, NRRL1951, obtained during WWII from an infected cantaloupe³. Biochemical and genetic analysis of industrial strains led to the identification of several important mutations in high-producing strains, including amplification of penicillin biosynthesis genes⁴. However, much of the molecular basis for improved productivity remains to be elucidated. A detailed understanding of the molecular biology of *P. chrysogenum* is not only relevant for 'classical' penicillins. By applying genetic engineering approaches, it has become possible to extend the range of fermentation products to include β -lactam

derivatives that could hitherto only be produced by chemical modification leading to great potential in terms of economy and sustainability. This is exemplified by the expression of the *Streptomyces clavuligerus* *cefE* gene, which encodes an expandase and has enabled high-yield production of cephalosporins with engineered *P. chrysogenum* strains⁵.

Accessibility to the full range of genomics techniques will be invaluable for further innovation in antibiotics production. Here, we present the complete genome sequence of *P. chrysogenum* Wisconsin54-1255 (ref. 6). An *in silico* analysis of the genome sequence has focused on key processes in penicillin production. Moreover, DNA microarrays have been applied for transcriptome comparisons of the sequenced strain and a derived high-producing strain.

RESULTS

Genome sequence and analysis

The *P. chrysogenum* genome was sequenced by the whole-genome sequencing method. The nuclear genome of 32.19 Mb was covered by 49 supercontigs, including 21 supercontigs larger than 5 kb and

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Table 1 Genome statistics overview

Nuclear genome	
General information	
Size (Mb)	32.2
G+C content (mole%)	48.9
Coding (%)	56.6
Gene number (with similarities)	12,943 (11,472)
Putative pseudogenes	592
Truncated ORFs	116
Questionable ORFs	2
Genes (<100 aminoacids)	283
Mean gene length (bp)	1,515
Genes with intron (%)	10,812 (83.5)
Exons	
Mean number per gene	3
Mean length (bp)	434
GC content (mole%)	52.8
Introns	
Mean number per gene	2.2
Mean length (bp)	87.4
GC content (mole%)	45.3
Intergenic regions	
Mean length (bp)	842
GC content (mole%)	44.4
RNA	
tRNA number	145
5S rRNA number	28
Mitochondrial genome	
Size (bp)	31,790
GC content (mole%)	24.9
Gene number	17
Mean gene length	985.4
Coding (%)	52.7
Genes with intron (%)	0
tRNA number	26

14 supercontigs larger than 100 kb. Annotation based on a minimum open reading frame (ORF) size of 100 amino acids revealed 13,653 ORFs (Table 1), including 592 probable pseudogenes and 116 ORFs whose sequences were truncated because their coding regions spanned contig borders. Two ORFs were considered unlikely to encode proteins because of their small size, absence of detectable protein motifs and low codon adaptation index. The sequenced mitochondrial genome comprised 31,790 bp and 17 identified ORFs.

BLASTP matches were found for 11,472 ORFs ($P < 0.001$) to a nonredundant

protein database, whereas the remaining 2,198 ORFs showed no significant similarities. Predicted protein-coding sequences account for 56.6% of the *P. chrysogenum* genome, with an average gene length of 1,515 bp. The GC content was 48.9% (52.8% for exons, 45.3% for introns and 44.4% for intergenic regions). On average, each gene contained 3.0 exons, with 83.5% of the genes containing introns. Using the FunCat classification system⁷, 5,329 of the 12,943 predicted nuclear-encoded proteins could be assigned to the functional protein classes metabolism, energy, cellular transport and protein fate (Fig. 1).

Comparison with other fungal genomes

The sequenced *P. chrysogenum* genome is comparable in size to that of other filamentous fungi (Supplementary Table 1 online). FunCat classification revealed a conserved orthologous core fungal proteome (Supplementary Fig. 1 online) involved in energy production, protein fate and cell fate. Phylogenetic analysis based on the concatenated protein set (Fig. 2a) confirmed a close relationship to *Aspergillus* species. The tree topology indicated that *P. chrysogenum* is only distantly related to the other two sequenced *Penicillium* species, *Penicillium marneffe* and *Talaromyces stipitatus* (teleomorph of *Penicillium stipitatus*). This contradicts a previously published phylogeny⁸ but is consistent with morphological observations⁹.

The 14 largest supercontigs (between 167 kb and 6,387 kb), presumably correspond to chromosome arms or even entire chromosomes. Alignment against various *Aspergilli* chromosomes suggests extensive reshuffling has occurred after divergence of the *Aspergillus* and *Penicillium* lineages (Fig. 2b). Several supercontigs are bounded by areas with multiple synteny breaks, which may correspond to subtelomeric regions (Fig. 2b). Indications for subtelomeric instability have also been observed in *Aspergilli*¹⁰ and *Magnaporthe oryzae*¹¹. Five supercontigs (nos. 12, 16, 20, 21 and 22) contain gaps surrounded by larger syntenic blocks, which appear to be recombination cold spots. These gaps resemble putative centromeres in the eight *A. fumigatus* chromosomes¹². Due to their high repeat content, centromeres as well as ribosomal DNA (rDNA) repeats regions typically do not get assembled into supercontigs in fungal genomes^{12,13}. Coincidentally, the upper gap on supercontig no.16 is not surrounded by large syntenic blocks and is likely to contain the rDNA region (Fig. 2b).

Genome alignment revealed four supercontigs (nos. 17, 19, 23 and 24), representing 4% of the genome, that show little similarity to

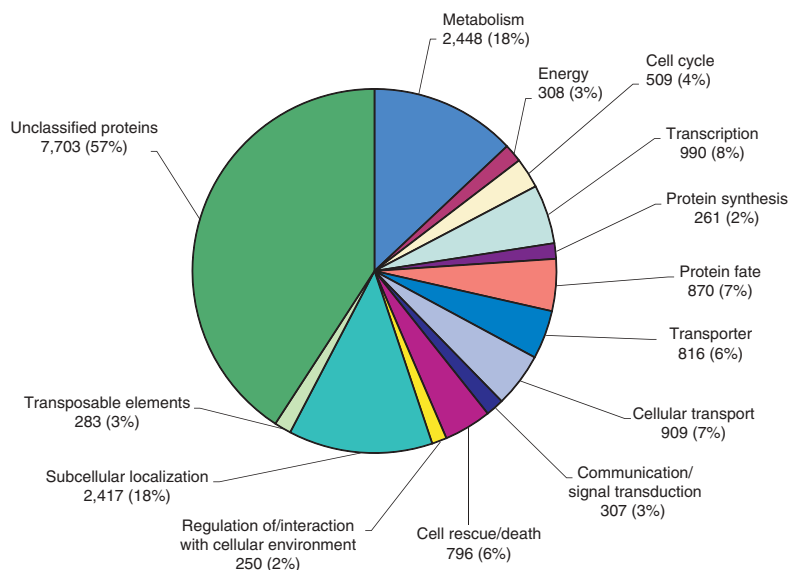
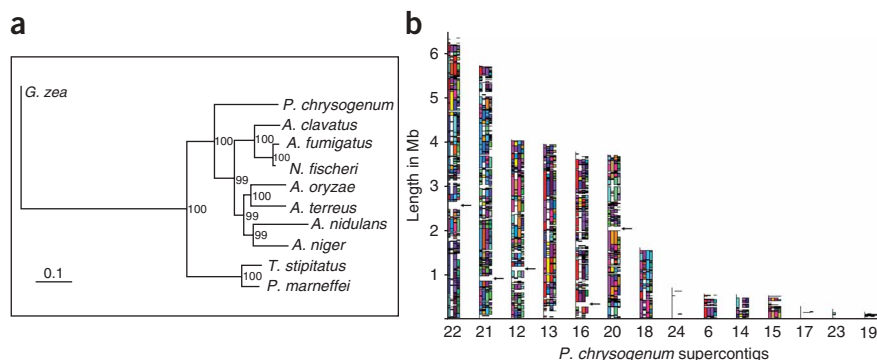


Figure 1 Genome characteristics of *P. chrysogenum*. Functional classification of *P. chrysogenum* ORFs. FunCat classes indicated are, 01 metabolism, 02 energy, 03 cell cycle and DNA processing, 04 transcription, 05 protein synthesis, 06 protein fate (folding, modification, destination), 07 transport facilitation, 08 cellular transport and transport mechanisms, 09 cellular communication/signal transduction mechanism, 10 cell rescue, defense and virulence, 11 regulation of/interaction with cellular environment, 12 subcellular localization, 13 transposable elements, 99 unclassified proteins.

Figure 2 Comparison of the *P. chrysogenum* genome to other sequenced filamentous fungi.

(a) Phylogenetic tree showing the relationships among sequenced *Aspergillus* and *Penicillium* species using *Gibberella zeae* as an outgroup. Branch lengths correspond to substitutions per site calculated using a maximum likelihood approach. Identical topologies were predicted using maximum parsimony and neighbor-joining methods. (b) Alignment of *A. nidulans*, *A. niger*, *A. fumigatus* and *A. oryzae* chromosomes against *P. chrysogenum* assemblies. Large assembled supercontigs (> 100 kb) from four *Aspergillus* genomes were aligned to the 14 largest *P. chrysogenum* assemblies using MUMmer

(<http://mummer.sourceforge.net/>). Regions with conserved gene order are represented by vertical columns of colored blocks. From left to right, *A. nidulans*, *A. niger*, *A. fumigatus* and *A. oryzae*, respectively. Each assembly (supercontig) from the target genomes is represented by a single color. Arrows indicate putative centromeres. Note: because the number of *A. nidulans* supercontigs (221) far exceeds the number of supercontigs available for the other *Aspergilli*, the length of aligned blocks between *A. nidulans* and *A. niger* may not represent the true extent of synteny between these two species.



Aspergilli. These regions contain *P. chrysogenum*-specific genes, which are typically smaller and contain fewer introns than other genes. Their biological roles are mostly unknown, although some seem to function in transport, metabolism or transcriptional regulation (Supplementary Fig. 2c online). These four nonsyntenic regions also contain numerous repeat elements and 23% of the genome's transposable elements (Supplementary Table 2 online). Similar genomic islands have been found in other fungal genomes^{11,13,14}.

Almost 30% of the predicted *P. chrysogenum* proteins lack orthologs in other sequenced fungi. In the closely related genus *Aspergillus*, the origin of lineage-specific genes has been largely attributed to either gene acquisition through horizontal gene transfer^{10,15} or to gene duplication followed by accelerated diversification and differential gene loss¹³. These genes tend to function in secondary metabolism and other accessory roles (Supplementary Fig. 2a) and may have a recent evolutionary origin. Phylogenetic analysis was applied to a subset of putative secondary metabolism genes. Thirty-three of such genes were identified using SMURF software (<http://www.tigr.org/software/>) encoding for: 20 polyketide synthases (PKS), 10 nonribosomal peptide synthetases (NRPS), 2 hybrid NRPS-PKS enzymes and 1 dimethylallyltryptophan synthase (Supplementary Table 3 online). This is similar to the numbers found in *Aspergilli*^{10,12,15,16}. The penicillin cluster is well known¹⁷, and the siderophore synthetases for ferri-chrome (Pc13g05250) and triacetylfulsarinine (Pc16g03850, Pc22g20400) were readily assigned by homology (Supplementary Table 4 online). None of the remaining six NRPS could be confidently identified. Pc21g15480 may encode roquefortine synthetase¹⁸ and is clustered with tryptophan dimethylallyl transferase (Pc21g15430). The putative tetrapeptide synthetases Pc13g14330 and Pc16g04690 have similar architectures to those in *Aspergilli* and may form cyclopeptides with two adjacent D-amino acids presumably related to malformin¹⁹. Pc21g10790 may form a cyclohexapeptide containing a fatty-acid derived component and is orthologous to a similar NRPS found in *A. oryzae*.

Penicillin biosynthetic genes

Several prokaryotic features of two penicillin biosynthetic genes, *pcbAB* and *pcbC*, encoding α -aminoacyl-tRNA synthetase and isopenicillinN (IPN) synthase, suggested that the penicillin gene cluster emerged through horizontal gene transfer from bacteria to fungi²⁰. Both genes lack introns (which is unique for large NRPS genes like *pcbAB*), are highly homologous to their bacterial

counterparts and are physically linked. Other features to consider are GC content (which is above 60% in prokaryotic penicillin producers) and specific codon usage. In *P. chrysogenum*, the GC content of the penicillin biosynthetic genes is only slightly higher than the overall genome average (Supplementary Data online). In the clavulanic acid producer *S. clavuligerus* the phenylalanine-codon UUU is extremely rare compared to UUG; UUU comprises only ~2.3% of total phenylalanine-codons. Whereas, in *P. chrysogenum*, the UUU codon overall is used in one-third of the cases, it is used for 26.2% and 17.6% of the phenylalanine-codons in *pcbAB* and *pcbC*, respectively. This can be interpreted as near complete codon adaptation because of the hypothesized transfer acquisition event.

Three other examples of possible horizontal gene transfer were identified in the *P. chrysogenum* genome: the arsenate-resistance cluster and two 6-methylsalicylic acid clusters (Supplementary Data and Supplementary Table 4). These gene clusters contain highly conserved bacterial-like genes with GC content well above the surrounding genes (exons with 55–58% GC).

The penicillin biosynthetic genes are clustered on supercontig 21 in the middle of a 120-kb region that is amplified in industrial *P. chrysogenum* strains⁴. Thirty-nine additional ORFs were identified in this region (Supplementary Table 5 online), including genes encoding transporters and transcriptional regulators. However, the predicted annotation of these ORFs does not suggest clear functions in penicillin biosynthesis as reported recently^{21,22}.

The third penicillin biosynthetic gene, *penDE*, encoding acyl-CoA: isopenicillinN acyltransferase, has a paralog, Pc13g09140. This gene was not transcribed under the conditions studied (Supplementary

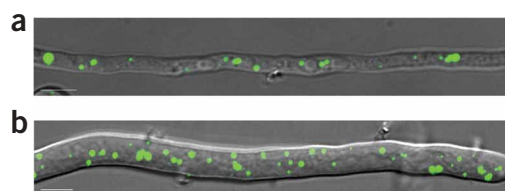


Figure 3 Microbodies in *P. chrysogenum* strains. (a,b) Cells of the *P. chrysogenum* type strain NRRL1951 (a) and the high penicillin producing strain DS17690 (b) producing the microbody-targeted protein GFP. SKL (Supplementary Data) were cultivated in batch cultures on penicillin-production media. Hyphae of strain DS17690 show enhanced microbodies numbers relative to the type strain NRRL1951. Scale bar, 5 μm.

Table 6 online). Detailed analyses are needed to reveal its actual function. Several orthologs of β -lactam biosynthesis genes were identified throughout the genome (**Supplementary Table 6**). As deletion of *phl*, which encodes phenylacetyl-CoA ligase, resulted only in a partial loss of penicillinG production²³, other phenylacetyl-CoA ligases must be present²⁴. These may include identified orthologs of 4-coumarate-CoA ligase. Surprisingly, several orthologs of bacterial²⁵ and fungal²⁶ isopenicillinN epimerase were identified. The predicted protein sequence of Pc12g11540 shares 40% homology with *S. clavuligerus* isopenicillinN epimerase, although it probably functions as an amino-transferase. Also, orthologs of *Acremonium chrysogenum* *cefD1* and *cefD2* were identified. The presence of these ORFs is remarkable, as *P. chrysogenum* can only produce penicillinN after introduction of both *A. chrysogenum* genes²⁷. The *P. chrysogenum* ORFs may be remnants of an ancestral cephalosporin pathway.

Microbodies

In *P. chrysogenum*, microbodies (peroxisomes) are essential for penicillin biosynthesis because the two final enzymatic steps catalyzed by acyl-CoA:isopenicillinN acyltransferase²⁸ and phenylacetyl-CoA ligase²⁹ are located in these organelles. Moreover, high-producing strains have enhanced microbody volume fractions (**Fig. 3**). Also, a further increase in microbody abundance by overexpression of the proliferation gene *pex11* leads to a significant increase in penicillin production³⁰. Genome 2D-searches³¹ with known consensus sequences for microbody targeting signals (PTS)³² identified 214 putative matrix proteins (196 and 17 with putative PTS1 and PTS2 respectively; 1 with both signals) (**Supplementary Table 7** online). Remarkably, the putative isopenicillinN-CoA epimerase (Pc22g13680) has a predicted PTS1. Many of the proteins are β -oxidation homologs, including multiple acyl-CoA synthetases and putative 3-ketoacyl-CoA

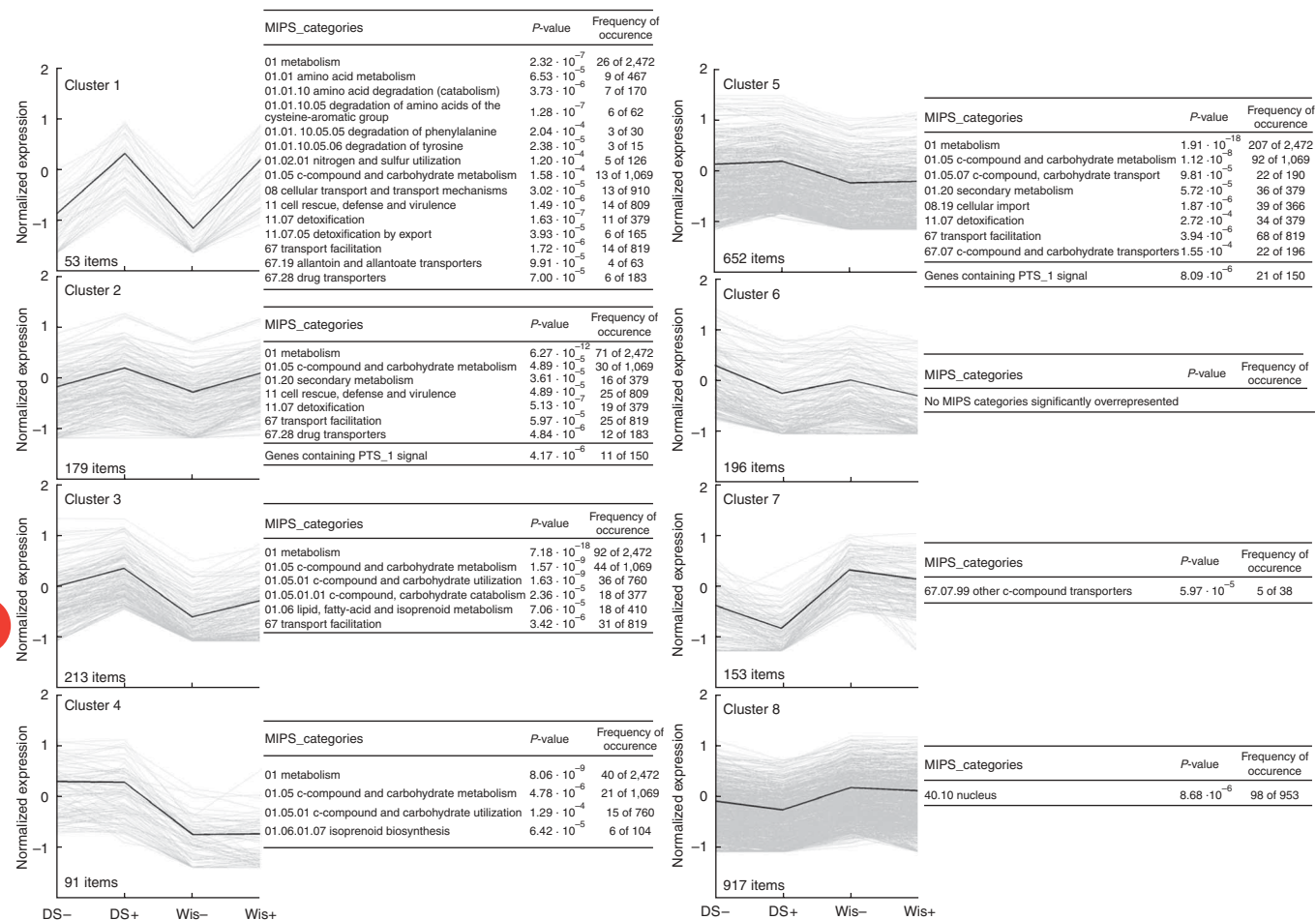


Figure 4 Transcriptional response of significantly changed genes, K-means clustered and the overrepresented functional categories in each cluster. Genes that showed a significantly different transcript level in at least one comparison (DS17690 + (DS+) versus – PAA (DS–); Wisconsin54-1255 + (Wis+) versus – PAA (Wis–); DS17690 + PAA versus Wisconsin54-1255 + PAA and DS17690 – PAA versus Wisconsin54-1255 – PAA) were grouped in eight clusters by K-means clustering. The thick lines represent the average of the mean normalized transcript levels of the genes in each cluster. The y axis represents $^{10}\log$ transcript levels. Three main categories were observed; (i) genes whose transcript levels were not influenced by the strain improvement program but with a higher transcript level in the presence of PAA (clusters 1 and 2); (ii) genes with a constitutively higher transcript level in DS17690 than in Wisconsin54-1255, irrespective of the presence of PAA (clusters 4 and 5); (iii) genes with a lower transcript level in DS17690, irrespective of the presence of PAA (clusters 7 and 8). Cluster 3 contains genes that showed a higher transcript level in DS17690 than in Wisconsin54-1255 and only responded to PAA in DS17690. Cluster 6 contains genes that only showed a lower transcript level in the presence of PAA in the DS17690 strain. Functional categories are mentioned together with a P-value indicating their overrepresentation in each cluster and the number of genes belonging to functional category in the cluster relative to the total number of these genes in the genome. Due to redundancy in the functional categories, statistically significant overrepresentation may not always reflect biological significance. Genes encoding enzymes containing the peroxisomal targeting signal PTS1 occurring in each cluster are indicated in the same manner.

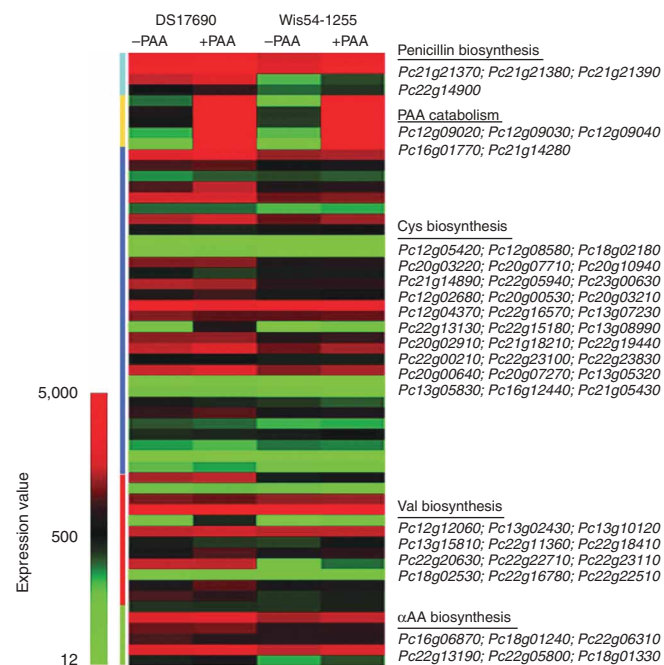


Figure 5 Eisen diagram of transcript levels of key genes in penicillin biosynthesis, amino-acid biosynthesis and phenylacetate catabolism. *P. chrysogenum* Wisconsin54-1255 and DS17690 strains were grown in glucose-limited chemostat cultures in the absence and presence of phenylacetic acid (PAA). The color bar indicates the range of the transcript levels for each gene, normalized to the average transcript level of the gene. Genes (putatively) related to penicillin biosynthesis, synthesis of the penicillin precursors cysteine, valine and α -aminoadipic acid and to phenylacetate catabolism are depicted.

thiolases (Pc13g12930, Pc15g00410 and Pc22g06820). Indeed, *P. chrysogenum* readily consumes oleate as the sole source of carbon and energy. Other PTS-containing proteins, such as D-amino acid oxidases, may play a role in the metabolism of various carbon or nitrogen sources.

Transcriptome analysis

When genomic DNA of *P. chrysogenum* was hybridized to microarrays, 99.4% of the probe sets hybridized (Supplementary Data). Once validated (Supplementary Data), these microarrays were used to investigate, at the transcriptome level, the molecular basis of the improved penicillin productivity achieved via classical strain improvement. Transcriptome analysis was performed on aerobic, glucose-limited chemostat cultures of Wisconsin54-1255 and the derived industrial, high-producing strain DS17690 (ref. 33). Some intermediates of β -lactam biosynthesis are produced in the absence of the side-chain phenylacetic acid (PAA)³³, but penicillinG biosynthesis is strictly dependent on PAA³⁴ (Supplementary Data). In DS17690 grown in the absence of PAA, 67% of the genome (~9,200 genes) yielded a detectable transcript (Supplementary Table 8 online). Under comparable conditions, 86% (~5500 genes) of the smaller *S. cerevisiae* genome was transcribed³⁵. To discriminate between PAA-responsive transcripts and transcripts potentially related to improved penicillinG productivity, the two strains were grown under penicillinG-producing and nonproducing conditions. In at least one of the four comparisons, 2,470 genes were differentially transcribed (Supplementary Table 8). By K-means clustering³⁶, these genes were assigned to eight clusters (Fig. 4 and Supplementary Tables 9–16 online).

Transcription of the penicillinG biosynthesis genes *pcbAB*, *pcbC* and *pil* was independent of PAA, but two- to fourfold higher in the high-producing strain (Fig. 4, cluster 5). *penDE* showed a similar trend (1.9- and 1.5-fold difference in the presence and absence of PAA, respectively; $P < 0.05$ in a *t*-test). Genes encoding enzymes involved in the biosynthesis of the amino-acid precursors of penicillin (cysteine, valine and α -aminoadipic acid) were also transcribed at higher levels in the high-producing strain independent of the presence of PAA (Fig. 4, cluster 5 and Fig. 5). This included sulfur reduction and early stages of serine (and cysteine) biosynthesis (Pc20g03220; Pc12g02680 and Pc12g04370), as well as a homolog of *O*-acetyl-homoserine (thiol)-lyase (Pc12g05420), a key enzyme in the trans-sulfuration pathway toward cysteine. Several genes encoding enzymes related to α -aminoadipic acid (lysine; Pc18g01330, Pc14g00150) and valine (Pc22g22510, Pc22g23110) metabolism showed a similar trend.

Of the genes predicted to encode microbody proteins, 27 showed higher transcript levels in DS17690, irrespective of PAA addition (Fig. 4, cluster 5). This class was also overrepresented among genes that were upregulated by addition of PAA (Fig. 4, cluster 2).

The homogentisate pathway for PAA degradation has been reported to be largely inactivated in Wisconsin54-1255 and, presumably, also in derived strains, owing to point mutations in the *pahA* gene encoding phenylacetate hydroxylase³⁷. Nevertheless, both strains showed very low, but significant rates of PAA consumption that could not be attributed to penicillinG production (Supplementary Data). Despite the low *in vivo* activity of this homogentisate pathway, its transcriptional regulation has been retained throughout the strain improvement program, as its structural genes showed increased transcript levels in the presence of PAA in both strains (Fig. 4, clusters 1 and 2).

Several transcriptional regulators have been implicated in the transcription of *pcbAB*, *pcbC* and *penDE* in *P. chrysogenum*³⁸. No penicillin-specific transcriptional regulator has been identified, although strong effects were reported from an enhancer sequence in the upstream region of *pcbAB*³⁹ and chromatin modulation⁴⁰. The *laeA* gene responsible for the latter effect strongly affects secondary metabolism in *Aspergillus fumigatus*⁴⁰. Although the *P. chrysogenum* ortholog was transcribed, its transcript levels were not substantially influenced by strain or cultivation conditions (Pc16g14010, Supplementary Table 17 online). Several other putative transcription factors, whose functions remain to be elucidated, were found to be associated with secondary metabolite clusters (Supplementary Data).

Transport

Transport mechanisms for β -lactam antibiotics and intermediates across the fungal plasma membrane and between intracellular compartments are poorly understood. Industrial fermentations yield high amounts of penicillinG in the external broth, whereas intracellular concentrations are typically tenfold lower. Moreover, penicillinG secretion is sensitive to verapamil⁴¹, an antagonist of multidrug transporters. This implies that secretion is an active process possibly mediated by (an) ABC transporter(s), whose identity has remained elusive. *P. chrysogenum* contains 830 genes that specify transporter proteins. Secondary transporters (688) are numerous with the majority belonging to the major facilitator superfamily (416), whereas 51 ABC transporters were identified. The functional categories metabolism, transport and detoxification were among the most strongly overrepresented in the gene clusters that were transcriptionally upregulated in the presence of PAA in both strains (Fig. 4, clusters 1 and 2 and Supplementary Table 18 online). Several of these showed

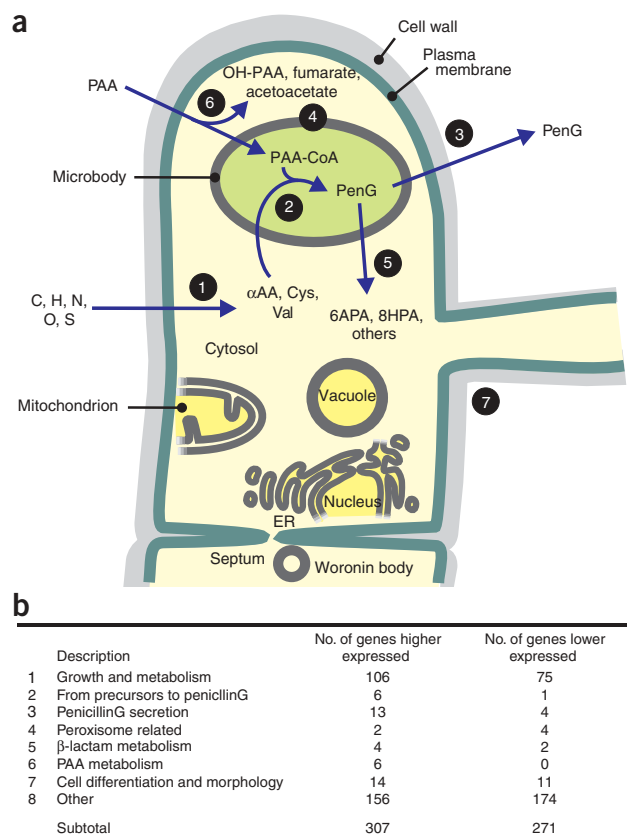


Figure 6 Categorization of differentially transcribed genes in DS17690 and Wisconsin54-1255. **(a)** Schematic representation of a *P. chrysogenum* cell and eight functional categories of cell metabolism directly influencing penicillin productivity: 1, growth and metabolism (from nutrients to precursors); 2, from precursors to penicillinG; 3, penicillinG secretion; 4, peroxisome related; 5, β-lactam metabolism; 6, PAA metabolism; 7, cell differentiation and morphology; 8, other (including unknowns, cell signaling and communication, transcription factors, stress response). **(b)** Transcript levels of all genes in DS17690 and Wisconsin54-1255 were compared under producing conditions (+PAA). Significant changes were selected by statistical analysis of microarrays (SAM) analysis with a threshold fold-change of 2 and a false-discovery rate of <1%, thus identifying 1,605 genes. An additional, more stringent criteria were imposed to select only genes whose highest average transcript level signal (in either strain) was larger than 200. The 578 remaining genes were functionally categorized according to the eight categories.

sequence similarity to known multidrug transporter genes. Transporters were significantly overrepresented within the class of genes expressed more highly in DS17690 than in Wisconsin54-1255, irrespective of PAA ($P = 3.94 \times 10^{-6}$, see Fig. 4, cluster 5), identifying sixty-eight potential active transporters (Supplementary Table 19 online). Interestingly, none of the previously suggested penicillin transporters^{41,42} are among this group. Although some transporter genes may be involved in transport of PAA rather than of β-lactams or intermediates, the strong enrichment of transporter genes suggest that penicillin secretion might result from the simultaneous activity of multiple transporters.

DISCUSSION

Although observations on amplification of penicillin biosynthetic genes⁴ and increased abundance of microbodies³⁰ have provided some insights, the question of how strain improvement transformed *P. chrysogenum* into an efficient penicillin producer has remained a formidable challenge. Availability of a complete genome sequence enables an integral approach. The potential is illustrated by several results, including increased transcript levels in a high-producing strain of genes involved in side-chain activation, of α-amino adipate, valine and cysteine biosynthesis genes, and of genes encoding multidrug transporters. Three hundred and seven of these upregulated genes were assigned to 8 categories directly influencing penicillin productivity (Fig. 6). A similar number of genes (271) showed reduced transcript levels in the high-producing strain (Fig. 6). Gene-by-gene analyses of this latter set suggests an overrepresentation of genes encoding morphogenesis and developmental factors. This is in line with the morphological differences between the two strains. These results indicate that metabolic control of penicillin production in high-producing strains is likely to be distributed over many cellular

processes that have been iteratively optimized in the long strain improvement history. Scattered gene duplications and deletions throughout the genome of different strains (Supplementary Data) may contribute to this combined up- and downregulation of different pathways and processes. However, most of these genes encode hypothetical proteins, except for Pc12g16490 (Supplementary Table 20 online, cluster no. 4), involved in cell development.

Despite the massive improvements already achieved in classical strain improvement, our results indicate that further improvement of penicillin production remains a possibility. For instance, the high-level expression of the PAA degradation pathway may impose an energetic burden and thus affect productivity. Moreover, the genome sequence shows evidence for other, potentially competing, β-lactam pathways that may contribute to by-product formation. Some of the genes, which strongly resemble genes encoding isopenicillinN epimerase and acetyltransferase, are highly transcribed and show increased levels in the high-producing strain DS17690. Further improvements can be anticipated from a detailed analysis and subsequent targeted modification of genes involved in microbody biogenesis and function, and of membrane transporters involved in the (intra)cellular transport of products and intermediates.

When a subset of the genes that showed altered transcript levels in DS17690 relative to Wisconsin54-1255 were knocked out, several amino-acid biosynthesis genes that were transcriptionally upregulated were confirmed to be involved in the increased β-lactam productivity (Supplementary Table 21 online). This is in contrast to the gene encoding Pc20g0240, a putative transporter with high mRNA levels in DS17690, which, when silenced, has little effect on β-lactam biosynthesis. Surprisingly, the knockout of a putative isopenicillinN-CoA epimerase seems to lead to a small increase in β-lactam productivity. In *Cephalosporium acremonium* this enzyme is involved in IPN to penicillinN conversion²⁷. It is tempting to speculate that this enzyme might be involved in a so far unidentified side reaction in *P. chrysogenum* reducing the overall yield of secreted penicillinG. These preliminary and diverse results demonstrate the need for a thorough follow-up of the transcriptome-based identification of targets for metabolic engineering.

The availability of a highly reproducible platform for transcriptome analysis will facilitate the systematic analysis of *P. chrysogenum* strain lineages, with the aim to further elucidate the molecular basis for the substantial increase in penicillinG productivity that has occurred during six decades of classical strain improvement. In addition, a full exploitation of the *P. chrysogenum* genome sequence will require the integration of additional levels of cellular information (for example, metabolome and proteome), as well as the construction of

genome-scale metabolic models⁴³. Such 'systems' approaches will ultimately contribute to further improvement of this important cell factory via inverse metabolic engineering⁴⁴.

METHODS

Genome sequencing and assembly. The genome of *P. chrysogenum* Wisconsin54-1255 (ATCC28089) was sequenced by the whole-genome random sequencing method¹² by obtaining paired-end reads from five libraries. Four plasmid libraries with inserts prepared by random shearing were constructed with insert sizes of 3–4 kb, 4–6 kb, 6–12 kb and 25–50 kb. Approximately 2.3× genome coverage was obtained from each of these libraries. In addition, 16,000 sequences were obtained from the ends of BAC clones. The BAC libraries were prepared by partial digestion of the *P. chrysogenum* genomic DNA using either *Bam*HI or *Hind*III⁴⁵. The mean insert size of these two libraries was 130 kb. Assembly of these sequences generated 49 supercontigs, with 14 of these larger than 100 kb. The sizes of the largest five supercontigs are 6.2 Mb, 5.6 Mb, 4.0 Mb, 3.9 Mb and 3.6 Mb. The final genome coverage in contigs was 9.8×. Genome assembly was accomplished using Celera Assembler⁴⁶. The mitochondrial sequence was obtained by manual review of the assembled contigs by sequence similarity to the mitochondrial sequence of *A. fumigatus* Af293. The genome sequence was deposited at EMBL under the accession numbers AM920416–a.m.920464.

Genome annotation and analysis. Analysis and annotation of the genomic sequences of *P. chrysogenum* was performed with a combined automatic and manual approach. For all supercontigs larger than 5 kb ORFs were predicted by a version of FGENESH⁴⁷ trained on sequences of Ascomycetes, as well as other algorithms (**Supplementary Methods** online). ORFs were named after the organism (Pc), supercontig number (two digits) followed by g (gene) and a five-digit number matching the order of the ORFs on the contig. For all ORFs identified from the above described approach an exhaustive automatic bioinformatic analysis in respect to function and structure of the respective protein was performed using the PEDANT-Pro^T software⁴⁸. Annotation of description, functional categories according to the Functional Catalog (FunCat) classification system⁷, and Enzyme Commission (EC) numbers have been performed for each *P. chrysogenum* ORF with a multi-step semiautomatic approach (**Supplementary Methods**). tRNA genes were identified as described in **Supplementary Methods**.

Repetitive elements. Identification of repeat elements was performed using RepeatMasker (<http://www.repeatmasker.org/>), RepeatScout (<http://repeatscout.bioprotejcts.org/>), Tandem Repeats Finder (<http://tandem.bu.edu/trf/trf.html>) and Transposon-PSI (Brian Haas, <http://transposonpsi.sourceforge.net>). The result of the latter was used to calculate the densities as the percentage of nucleotide bases in the regions of interest that overlap with repeat or transposable elements (**Supplementary Table 2**).

Genome-genome alignment. Pair-wise alignments between *P. chrysogenum* and four target genomes (*A. fumigatus*, *A. oryzae*, *A. niger* and *A. nidulans*) were performed using the Promer program of the MUMmer package (<http://mummer.sourceforge.net/>). Supercontigs larger than 100 kb of the four *Aspergillus* species were aligned against the 14 largest *P. chrysogenum* supercontigs.

Phylogenetic analysis and species tree. To generate the species tree, a total of 90 orthologous genes from *P. chrysogenum*, *P. marneffei* (GenBank ABAR000000000), *T. stipitatus* (GenBank ABAS000000000), *A. niger*¹⁶, *A. nidulans*¹⁰, *A. oryzae*¹⁵, *A. fumigatus*¹², *A. clavatus* (GenBank AAKD000000000), *A. terreus* (Refseq NT_165925–NT_165951), *Neosartorya fischeri* (GenBank AAKE000000000) and *Gibberella zeae* (anamorph *Fusarium graminearum*)⁴⁹ were aligned at the amino acid level (**Supplementary Table 22** online). To minimize the effect of incorrect or incongruent gene models, these proteins were chosen on the basis of having identical numbers of introns in each species and similar lengths (95% overlap). Sequences were aligned using Muscle⁵⁰. DNA alignments were then concatenated and passed to the Phylip package. Maximum likelihood trees were calculated on each replicate, applying the JTT

substitution model with a gamma distribution ($\alpha = 0.5$) of rates over four categories of variable sites, and a consensus tree was produced.

Chemostat cultivations. Independent triplicate chemostat cultures of *P. chrysogenum* Wisconsin54-1255 and the high-producing penicillinG strain DS17690 were run in the presence and absence of phenylacetic acid exactly as described before³³.

Microarray methods (detailed procedures are available in **Supplementary Methods**). The *P. chrysogenum* genome sequence information was used to prepare a proprietary DNA microarray, using the Affymetrix Custom GeneChip program (Affymetrix): GeneChip, DSM_PENa520255F. Samples from chemostat cultures were filtered within seconds and quenched in liquid nitrogen. Total RNA isolation with Trizol reagent and acid phenol-chloroform for extraction was followed by cDNA synthesis and cRNA synthesis. Hybridized arrays were scanned and analyzed using the Affymetrix GeneChip Operating Software (GCOS, Affymetrix). All arrays were globally scaled to a target value of 100 using the average signal from all gene features using GCOS. Differential expression was assessed using the Significance Analysis of Micro arrays (SAM version 1.21) add-in to Microsoft Excel³⁴. The fold-change threshold and the false discovery rate values were set at 2% and 1%, respectively. Enrichment of Munich Information Center for Protein Sequences (MIPS) categories was assessed by Fisher's Exact test employing hypergeometric distribution with a *P*-value cut-off of $3 \cdot 10^{-4}$ (with a Bonferroni correction). The genes with significantly changed expression in one of the comparisons were arranged in clusters by the *K*-means clustering tool of Genedata Expressionist (Genedata). The coefficient of variation (CV) of transcript levels of independent, triplicate chemostat cultures did not exceed 21% (**Supplementary Data**) and was similar to that of transcriptome analysis on chemostat cultures of the nonfilamentous fungus *S. cerevisiae*³⁴.

Accession number. The array data were deposited at Genome Expression omnibus under the serial number GSE9825. The genome sequence was deposited at EMBL under the accession numbers AM920416–AM920464.

Note: Supplementary information is available on the Nature Biotechnology website.

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AUTHOR CONTRIBUTIONS

M.A.v.d.B. initiated and coordinated the *P. chrysogenum* genome project. W.C.N. coordinated the genome sequencing. N.D.F., J.H.B., V.J. and J.W. were responsible for genome assembly and genome-genome comparisons. K.A., R.A. and C.W. were responsible for genome annotation and analysis. C.G.-E. and J.E.M. performed the analysis of transcription factors. H.v.d. and N.D.F. analysed the secondary metabolites and gene clusters. J.A.K.W.K., I.J.v.d.K. and M.V. performed the analysis of the microbody proteins and induction. A.K., J.G.N. and A.J.M.D. carried out the transporter analysis. N.N.M.E.v.P. was responsible for the microarray design in collaboration with Affymetrix. W.H.M.H. and J.A.R. performed MicroArray data analyses. D.M.H., J.T.P. and

J.-M.D. were responsible for chemostat cultivations and subsequent MicroArray data analyses. J.G.N. performed the analysis of MFS and A.K. did the analysis of ABC transporter clusters. J.T.P., A.J.M.D., W.C.N., R.A.L.B. and M.A.v.d.B. wrote the final text of the manuscript.

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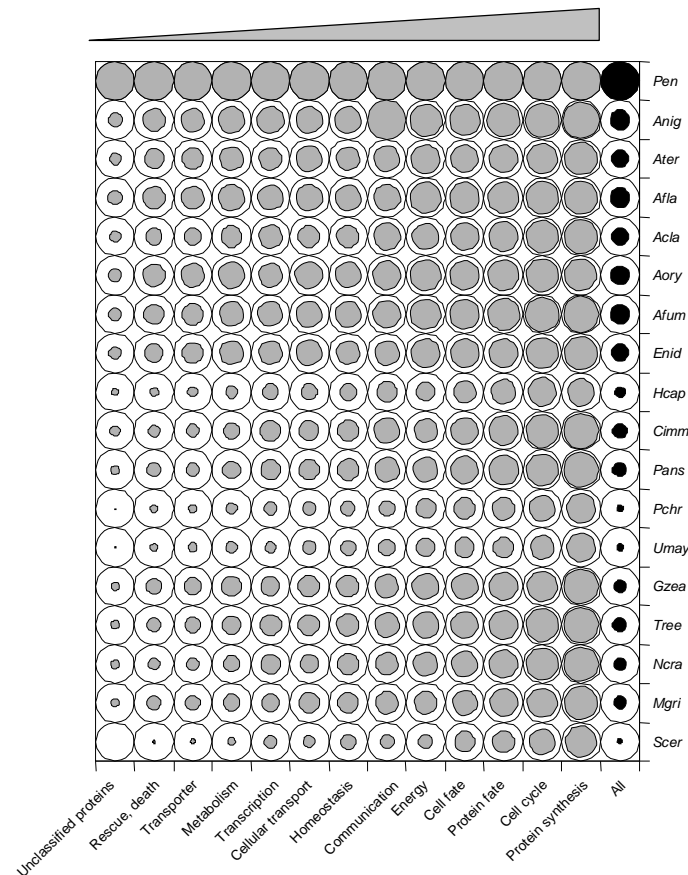
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Supplementary Figure 1. Ortholog comparison of main functional classes in different fungal genomes. Pen: *Penicillium*

chrysogenum, Anig: *Aspergillus niger*, Ater: *Aspergillus terreus*, Afla: *Aspergillus flavus*, Acla: *Aspergillus clavatus*, Aory: *Aspergillus oryzae*, Afum: *Aspergillus fumigatus*, Enid: *Emericella nidulans*, Hcap: *Histoplasma capsulatum*, Cimm: *Coccidioides immitis*, Pans: *Podospora anserina*, Pchr: *Phanerochaete chrysosporium*, Umay: *Ustilago maydis*, Gzea: *Gibberella zeae*, Tree: *Trichoderma reesei*, Ncra: *Neurospora crassa*, Mgri: *Magnaporthe grisea*, Scer: *Saccharomyces cerevisiae*.

The size of the filled circles is proportional to the number of ortholog genes in each category. Categories have been sorted in respect to the number of ortholog genes over all indicated genomes increasing from left to right. Category “All” represents the total number of ortholog genes in respect to *P. chrysogenum* for the listed fungal genomes.



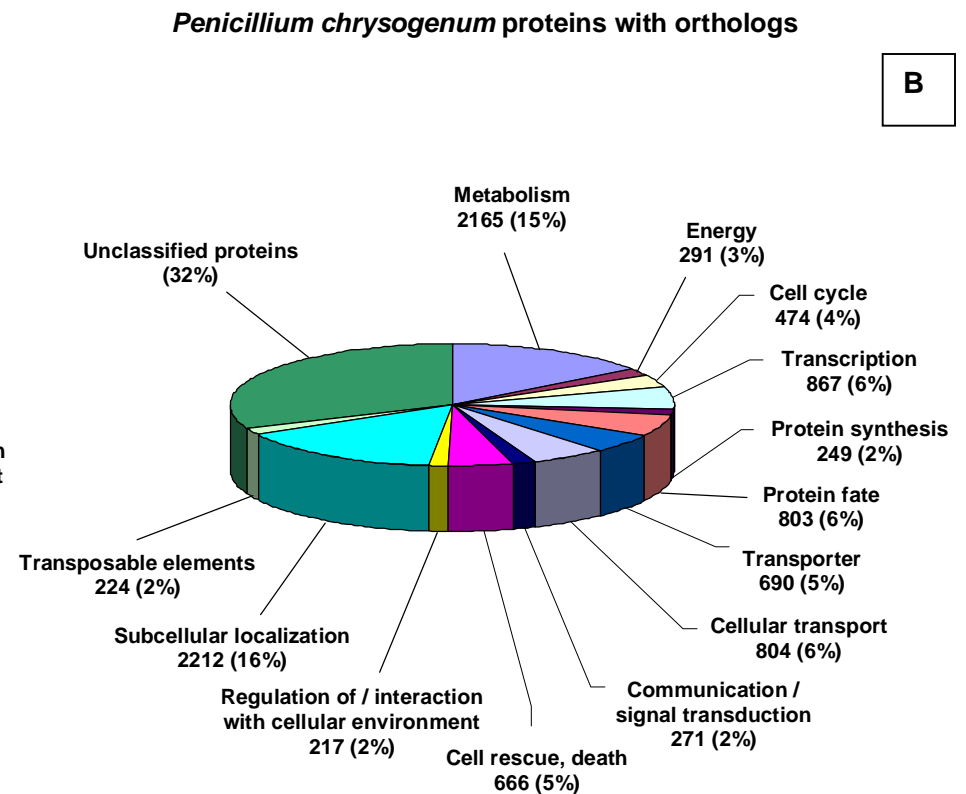
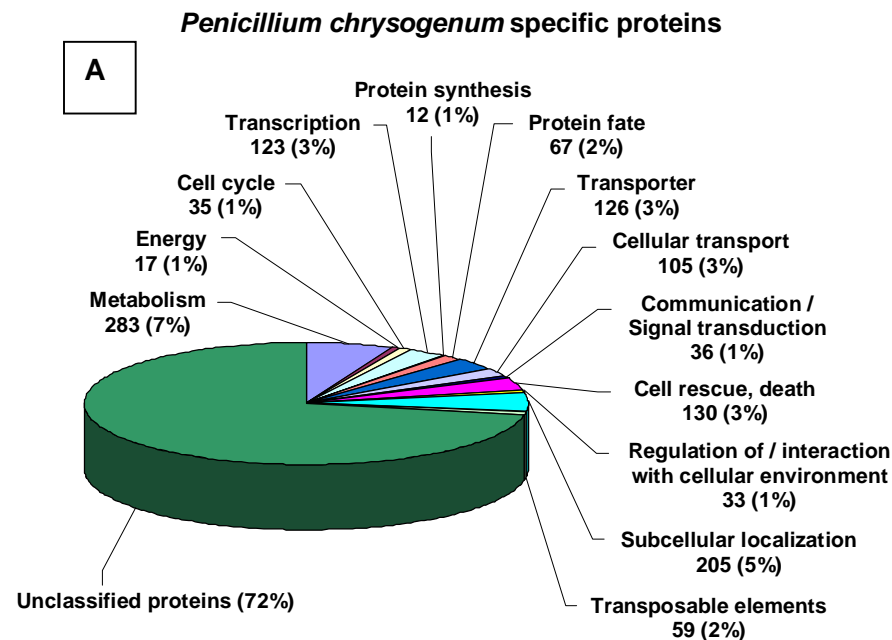
Supplementary Figure 2. Ortholog comparison of functional classes related to metabolism and energy in different fungal genomes. Pen: *Penicillium chrysogenum*, Anig: *Aspergillus niger*, Ater: *Aspergillus terreus*, Afla: *Aspergillus flavus*, Acla: *Aspergillus clavatus*, Aory: *Aspergillus oryzae*, Afum: *Aspergillus fumigatus*, Enid: *Emericella nidulans*, Hcap: *Histoplasma capsulatum*, Cimm: *Coccidioides immitis*, Pans: *Podospira anserina*, Pchr: *Phanerochaete chrysosporium*, Umay: *Ustilago maydis*, Gzea: *Gibberella zeae*, Tree: *Trichoderma reesei*, Ncra: *Neurospora crassa*, Mgri: *Magnaporthe grisea*, Scer: *Saccharomyces cerevisiae*.

The size of the filled circles is proportional to the number of ortholog genes in each category. Categories have been sorted in respect to the number of ortholog genes over all indicated genomes increasing from left to right.

The numbers on the x-Axis represent the following functional categories:

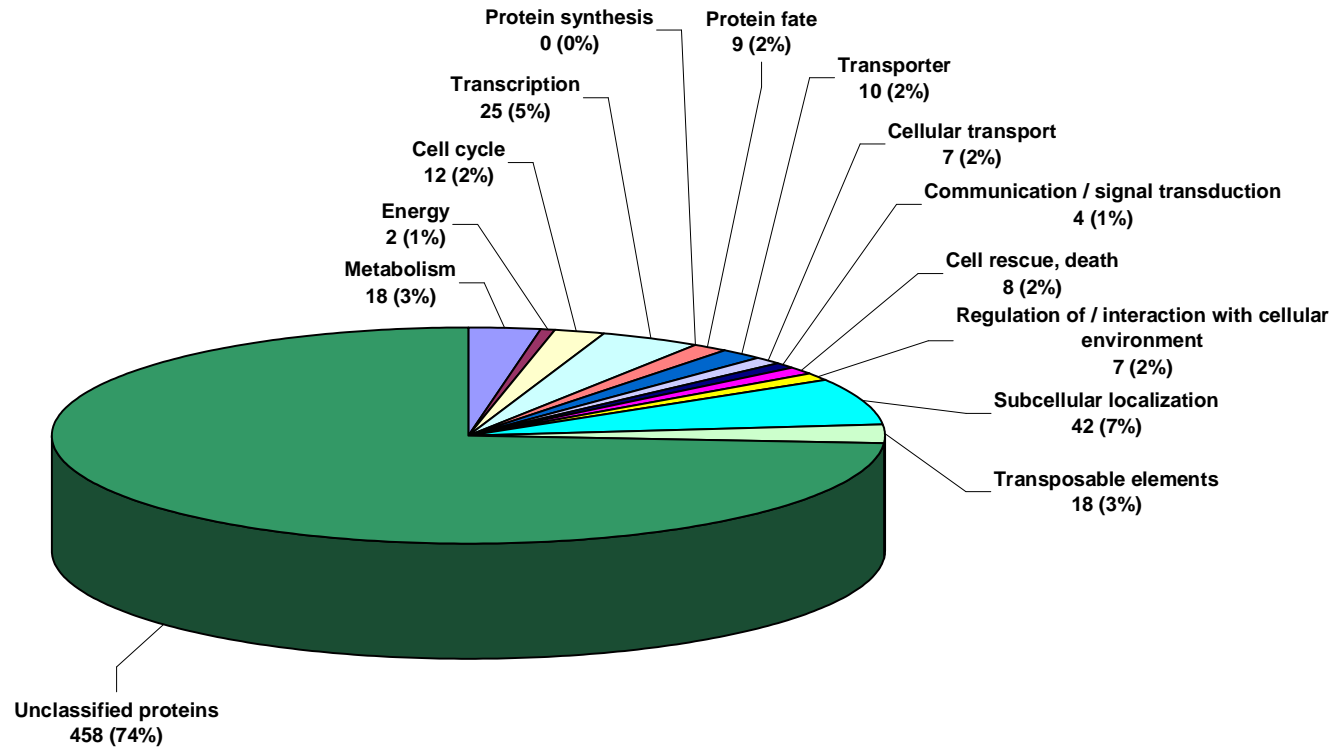
1 biosynthesis of nonprotein amino acids, **2** biosynthesis of polyketides, **3** biosynthesis of alkanes, alkenes, alkanals, alkanols, **4** anaerobic aromate catabolism, **5** biosynthesis of aminoglycoside antibiotics, **6** biosynthesis of β -lactams, **7** biosynthesis of peptide antibiotics, **8** aliphatic hydrocarbon catabolism, **9** biosynthesis of secondary products derived from L-phenylalanine and L-tyrosine, **10** isoprenoid biosynthesis, **11** aerobic aromate catabolism, **12** catabolism of secondary metabolites, **13** biosynthesis of amines, **14** fermentation, **15** aminosaccharide biosynthesis, **16** biosynthesis of alkaloids, **17** nitrogen and sulfur metabolism, **18** degradation of amino acids of the cysteine-aromatic group, **19** biosynthesis of acetoacetate, acetone, hydroxybutyric acid, **20** degradation of amino acids of the glutamate group, **21** assimilation of ammonia, biosynthesis of the glutamate group, **22** breakdown of lipids, fatty acids and isoprenoids, **23** degradation of amino acids of the aspartate group, **24** pentose-phosphate pathway oxidative branch, **25** metabolism of energy reserves (e.g. glycogen, trehalose), **26** fatty acid biosynthesis **27** biosynthesis of derivatives of dehydroquinic acid, shikimic acid and chorismic acid, **28** purine nucleotide metabolism, **29** degradation of amino acids of the pyruvate family, **30** glycolipid biosynthesis, **31** phosphate metabolism, **32** oxidation of fatty acids, **33** polysaccharide biosynthesis, **34** biosynthesis of secondary monosaccharides, **35** biosynthesis of secondary products derived from L-lysine, L-arginine and L-histidine, **36** extracellular metabolism, **37** biosynthesis of the cysteine-aromatic group, **38** glyoxylate cycle, **39** biosynthesis of the aspartate family, **40** glycolysis and gluconeogenesis, **41** metabolism of vitamins, cofactors, and prosthetic groups, **42** polynucleotide degradation, **43** deoxyribonucleotide metabolism, **44** pyrimidine nucleotide metabolism, **45** biosynthesis of the pyruvate family (alanine, isoleucine,

leucine, valine) and D-alanine, **46** biosynthesis of secondary products derived from L-glutamic acid, L-proline and L-ornithine, **47** urea cycle, biosynthesis of polyamines and creatine, **48** biosynthesis of secondary products derived from L-tryptophan, **49** electron transport and membrane-associated energy conservation, **50** tricarboxylic-acid pathway (citrate cycle, Krebs cycle, TCA cycle), **51** respiration, **52** phospholipid biosynthesis, **53** C-1 compound catabolism, **54** biosynthesis of porphyrins, **55** biosynthesis of glycosides, **56** biosynthesis of sulfuric acid and L-cysteine derivatives, **57** metabolism of cyclic and unusual nucleotides, **58** pentose-phosphate pathway non oxidative branch, **59** degradation of amino acids of the hydroxyamino-acid group, **60** biosynthesis of derivatives of homoisopentenyl pyrophosphate, **61** biosynthesis of cobalamins.



C

all 4 non-syntenic assemblies



Supplementary Table 1. Genome statistics comparison between different filamentous fungi.

	<i>P. chrysogenum</i>	<i>A.niger</i>	<i>A. nidulans</i>	<i>A. fumigatus</i> Af293
Chromosomes/Scaffolds				
Total Length (Kb)	32,224	33,931	30,069	28,810
GC content (%)	48.9	50.4	50.0	49.8
Number of Protein Coding Genes	12,941	14,165	10,662	9,632
Mean Gene Length (bp)	1,515	1,573	1,868	1,478.1
Gene Density	2,490	2,395	3,151	2,990
Percent Coding (%)	56.6	55.2	50.0	49.4
Genes with Introns (%)	83.5	87.0	86.9	78.6
Exons				
Average size (bp)	434	370	436	504
Number	41,996	50,629	35,797	28,254
Mean # per Gene	3	3.5	3.4	2.8
GC Content (%)	52.9	53.7	53.3	54
Total Length (bp)	18,238,634	18,733,984	15,477,748	14,241,720
Introns				
Average size (bp)	87.4	97.2	91	82
Number	28,326	36,464	24,792	18,619
Mean # per Gene	2.2	2.6	2.4	1.8
GC Content (%)	45.3	45.3	45.8	46.7
Total Length (bp)	2,475,921	3,544,638	2,243,391	1,521,138
Intergenic Region				
GC Content (%)	44.4	46.4	47.4	45.9
Mean Length (bp)	842	822	1,137	1,322
Longest intergenic region (bp)	44,111	19,212	13,654	55,846
RNA				
tRNA number	145	269	n/a	179
5S rRNA number	28	56	n/a	33

Supplementary table 2. Features of the four non-syntenic supercontigs

Non-syntenic Contigs	67	69	73	74	All four	Genome
Length (Kb)	292	168	229	709	1398	32183
Coding (%)	33.8	37.5	36.3	35.0	35.3	56.6
C+G content (%)	48.6	48.1	48.0	48.1	48.2	48.9
Genes	128	78	103	327	636	13653
Genes (< 100 aa)	6	3	4	8	21	283
Genes with introns (%)	71.1	70.5	82.5	76.1	75.5	83.5
Mean No. exons	2.78	2.50	3.07	2.67	2.75	3.07
Mean protein length (bp)	302	309	307	288	302	446
Mean gene length (aa)	905	927	922	865	905	1338
Repeat element density	0.0177	0.0163	0.0196	0.0158	0.01735	0.0104
TE elements	30	25	11	65	131	559
Pseudogenes	19	11	13	40	83	592

Supplementary Table 3. Detailed analysis of putative key-enzymes in secondary metabolite synthesis

Analysis was performed using SMURF software (<http://www.tigr.org/software/>)

Acyl_transf, acyl transferase domain; AMP, adenosyl-monophosphate domain; C, condensation domain; BKS-C, domain with ubiquitin ligase activity, C-terminal; BKS-N, domain with ubiquitin ligase activity, N-terminal; PP, pyrophosphate domain; NAD_4, NAD-binding domain; adh_short, short chain alcohol dehydrogenase domain; DMAT, dimethylallyltransferase domain; arom_pren_DMAT, dimethylallyltransferase domain involved in prenylation of aromatic ring systems; TDMAT, tryptophan-DMAT domain; Lys2, alpha aminoadipate reductase domain; alpha_am_amid - alpha aminoadipate domain

orf code	orf description	Proposed enzyme type	Domains												
			Acyl_transf	AMP	C	BKS-C	BKS-N	PP	NAD_4	adh_short	DMAT	arom_pren_DMAT	TDMAT	Lys2	alpha_am_amid
Pc21g15430	dimethylallyltryptophan synthase dmaW	DMAT	0	0	0	0	0	0	0	0	1	1	0	0	0
Pc14g00080	hybrid NRPS/PKS enzyme, putative	HYBRID	1	1	1	1	1	2	1	1	0	0	0	1	1
Pc16g13930	hybrid NRPS/PKS enzyme, putative	HYBRID	1	1	1	1	1	1	1	1	0	0	0	1	1
Pc13g05250	cyclic peptide AM-toxin synthase like protein An06g01300	NRPS	0	3	6	0	0	6	0	0	0	0	0	0	1
Pc13g14330	HC-toxin peptide synthase HTS	NRPS	0	4	6	0	0	4	0	0	0	0	0	0	1
Pc16g03850	peptide synthase pesA	NRPS	0	1	2	0	0	2	0	0	0	0	0	0	1
Pc16g04690	peptide synthase pesA	NRPS	0	4	6	0	0	5	0	0	0	0	0	0	1
Pc21g01710	peptide synthase like protein An04g06260	NRPS	0	2	1	0	0	2	0	0	0	0	0	0	1
Pc21g10790	HC-toxin peptide synthase HTS	NRPS	0	6	7	0	0	6	0	0	0	0	0	0	1
Pc21g12630	peptide synthase pesA	NRPS	0	2	2	0	0	2	0	0	0	0	0	0	1
Pc21g15480	cyclic peptide AM-toxin synthase like protein An03g00650	NRPS	0	2	2	0	0	2	0	0	0	0	0	0	1
Pc21g21390	nonribosomal alpha-aminoadipyl-cysteinyl-valine synthase PcbAB/AcvA	NRPS	0	3	3	0	0	3	0	0	0	0	0	0	1
Pc22g00400	cyclic peptide AM-toxin synthase AMT	NRPS	0	1	2	0	0	2	0	0	0	0	0	0	1
Pc06g01540	saframycin Mx1 synthase safA	NRPS-Like	0	1	0	0	0	1	1	0	0	0	0	1	0
Pc12g09980	AMP-binding enzyme, putative	NRPS-Like	0	1	0	0	0	1	0	0	0	0	0	0	0
Pc12g13170	polyketide synthase PKS17	NRPS-Like	0	1	0	0	0	1	1	0	0	0	0	1	1
Pc13g12570	saframycin Mx1 synthase safA	NRPS-Like	0	1	0	0	0	1	1	0	0	0	0	1	1
Pc14g01790	polyketide synthase PKS17	NRPS-Like	0	1	0	0	0	0	1	0	0	0	0	1	1
Pc16g09930	FKS20 biosynthetic gene cluster polyketide synthase like protein An12g02750	NRPS-Like	0	1	0	0	0	1	0	0	0	0	0	0	0
Pc18g00380	saframycin Mx1 synthase safA	NRPS-Like	0	1	0	0	0	1	1	1	0	0	0	1	1
Pc20g02260	aminoadipate reductase enzyme lys2	NRPS-Like	0	1	0	0	0	0	1	0	0	0	0	1	1
Pc20g02590	saframycin Mx1 synthase safA	NRPS-Like	0	1	0	0	0	1	1	0	0	0	0	1	0
Pc20g09690	polyketide synthase PKS17	NRPS-Like	0	1	0	0	0	1	1	0	0	0	0	1	1
Pc20g12670	hypothetical protein 1457_scaffold_8.tfa_130cg	NRPS-Like	0	1	0	0	0	0	1	0	0	0	0	1	1
Pc21g22530	hypothetical protein contig_1_135_scaffold_11.tfa_110wg	NRPS-Like	0	1	0	0	0	1	0	0	0	0	0	0	1
Pc21g22650	polyketide synthase PKS17	NRPS-Like	0	1	0	0	0	0	1	0	0	0	0	1	1
Pc22g06310	L-aminoadipate-semialdehyde dehydrogenase large subunit Lys2 - Penicillium chrysogenum	NRPS-Like	0	1	0	0	0	1	1	0	0	0	0	1	1
Pc22g09430	NRPS-like enzyme, putative	NRPS-Like	0	1	0	0	0	0	1	0	0	0	0	1	1
Pc12g05590	nonaketide synthase lovB	PKS	1	0	0	1	1	0	0	1	0	0	0	0	0
Pc13g04470	lovastatin diketide synthase lovF	PKS	1	0	0	1	1	0	0	1	0	0	0	0	0
Pc13g08690	lovastatin diketide synthase lovF	PKS	1	0	0	1	1	0	0	1	0	0	0	0	0
Pc16g00370	6-methylsalicylic acid synthase like protein An10g00140	PKS	1	0	0	1	1	1	0	1	0	0	0	0	0
Pc16g03800	polyketide synthase PKS1	PKS	1	0	0	1	1	0	0	1	0	0	0	0	0
Pc16g04890	polyketide synthase like protein An01g01130	PKS	1	0	0	1	1	0	0	1	0	0	0	0	0
Pc16g11480	lovastatin diketide synthase lovF	PKS	1	0	0	1	1	0	0	1	0	0	0	0	0
Pc21g00960	lovastatin diketide synthase lovF	PKS	1	0	0	1	1	0	1	1	0	0	0	1	1
Pc21g03930	lovastatin diketide synthase lovF	PKS	1	0	0	1	1	0	0	1	0	0	0	0	0
Pc21g03990	polyketide synthase like protein An15g05090	PKS	1	0	0	1	1	0	0	1	0	0	0	0	0
Pc21g04840	polyketide synthase PKS1	PKS	1	0	0	1	1	0	0	1	0	0	0	0	0
Pc21g05070	polyketide synthase like protein An09g01860	PKS	1	0	0	1	1	1	1	0	0	0	0	1	0
Pc21g05080	lovastatin diketide synthase lovF	PKS	1	0	0	1	1	0	0	1	0	0	0	0	0
Pc21g12440	lovastatin diketide synthase lovF	PKS	1	0	0	1	1	0	0	1	0	0	0	0	0
Pc21g12450	polyketide synthase wA	PKS	1	0	0	1	1	1	0	0	0	0	0	0	0
Pc21g15160	lovastatin diketide synthase lovF	PKS	1	0	0	1	1	0	0	1	0	0	0	0	0
Pc21g16000	polyketide synthase PKS1	PKS	1	0	0	1	1	2	0	0	0	0	0	0	0
Pc22g08170	6-methylsalicylic acid synthase like protein An10g00140	PKS	1	0	0	1	1	1	0	1	0	0	0	0	0
Pc22g22850	polyketide synthase like protein An09g01860	PKS	1	0	0	1	1	1	0	0	0	0	0	0	0
Pc22g23750	lovastatin diketide synthase lovF	PKS	1	0	0	1	1	0	0	1	0	0	0	0	0
Pc12g02940	lovastatin diketide synthase lovF	PKS-Like	1	0	0	1	1	0	0	0	0	0	0	0	0
Pc16g03760	strong similarity to polyketide synthase PKS1 - Cochliobolus heterostrophus [putative]	PKS-Like	0	0	0	0	0	0	0	1	0	0	0	0	0
Pc22g09030	3-oxoacyl-[acyl-carrier-protein]-synthase, putative	PKS-Like	0	0	0	1	1	0	0	0	0	0	0	0	0

Supplementary Table 4. PKS and NRPS clusters in *P. chrysogenum* genome

DMAT				
NRPS				
NRPS-Like				
PKS				
PKS-Like	Identified by SMURF (Supplementary Table 7)			
PKS-like	Not identified via SMURF			
Hybrid				

Secondary metabolite gene or cluster	ORF code	Description of putative <i>P. chrysogenum</i> ORF	Detailed analyses	Average transcript levels @			
				WIS -PAA	WIS +PAA	DS -PAA	DS +PAA
	Pc06g01490	strong similarity to hypothetical amine transporter SPCC18.02 - Schizosaccharomyces pombe	arabinose efflux permease, weakly similar protein in Ajellomyces capsulatus (35% identity, EDN06642)	57.4	41.2	14.2	0.4
	Pc06g01500	strong similarity to hypothetical protein An12g02740 - Aspergillus niger	Predicted protein with SAM methyltransferase domain, weakly similar proteins in A. terreus (35% identity, EAU36191) and A. niger (32% identity, An12g02740)	84.6	57.8	63.5	35.1
	Pc06g01510	similarity to hypothetical protein AAM35689.1 - Xanthomonas axonopodis	Hypothetical protein, similar proteins in Ajellomyces capsulatus (44% identity, EDN11135) and A. oryzae (43% identity, BAE63447)	55.3	36.4	25.7	9.3
	Pc06g01520	strong similarity to hypothetical protein 1323_scaffold_5.tfa_400cg - Fusarium graminearum	Putative benzoate 4-monoxygenase cytochrome P450, related enzymes in G Zeae (41% identity, XP38819), A. clavatus (39% identity, EAW14333), N. fischeri (39% identity, EAW18955), A. oryzae (39% identity, BAE65144)	52.6	31.0	19.0	5.6
	Pc06g01530	similarity to hypothetical protein 4 - Streptomyces coelicolor	NmrA-family, negative transcriptional regulator, related proteins in Ajellomyces capsulatus	33.9	24.4	56.0	3.0
1	Pc06g01540	strong similarity to saframycin Mx1 synthase safA - Myxococcus xanthus	Monomodular NRPS-like enzyme, 1025 AA, Highly similar NRPS in A. oryzae (64% identity, gi 83773330 dbj BAE63457.1), N. fischeri (55% identity, gi 119415803 gb EAW25740.1), and Ajellomyces capsulatus (52% identity, gi 150415793 gb EDN11137.1)	11.2	5.4	15.6	1.7
	Pc12g02900	strong similarity to transposase Tan1 - Aspergillus niger [putative pseudogene]		135.0	86.7	72.1	26.9
	Pc12g02910	hypothetical protein		983.7	1044.6	575.7	240.9
	Pc12g02920	weak similarity to 2-hydroxyisoflavone reductase IRL - Zea mays		126.3	58.5	152.4	152.5
	Pc12g02930	strong similarity to oxidoreductase like protein An09g01260 - Aspergillus niger		2.4	1.3	24.5	4.7
	Pc12g02940	strong similarity to lovastatin diketide synthase lovF - Aspergillus terreus [putative sequencing error]	PKS-like enzyme, 2523 AA, 13 introns, Related PKS in A. oryzae (45% identity, gi 83764443 dbj BAE54587.1 and gi 83765672 dbj BAE55815.1), N. fischeri (45% identity, gi 119406436 gb EAW16387.1), A. niger (44% identity, gi 119406436 gb EAW16387.1), A. oryzae (43% identity, gi 83773906 dbj BAE64031.1)	4.7	2.2	105.9	20.0
2	Pc12g02950	hypothetical protein		424.5	119.3	179.4	22.6
	Pc12g02960	strong similarity to hypothetical protein 1384_scaffold_6.tfa_150cg - Fusarium graminearum	Putative acetyltransferase, similar enzyme in A. terreus (56% identity, gb EAU33508.1)	18.9	24.6	82.8	77.8
3	Pc12g05590	strong similarity to nonaketide synthase lovB - Aspergillus terreus	polyketide synthase, related to equisetin synthase, 2715 AA, 10 introns, related enzymes e.g. in A.	3.9	2.9	4.8	6.2
	Pc12g05600	strong similarity to H+-biotin symporter Vht1 - Saccharomyces cerevisiae [putative sequencing error]		1.4	1.9	2.4	1.9
	Pc12g05610	strong similarity to hypothetical protein An01g10470 - Aspergillus niger [putative sequencing error]		69.2	61.8	45.3	47.0
4	Pc12g09980	Monomodular NRPS-like enzyme with additional transferase hexapeptide repeats, 1619 AA, 2	strong similarity to acid-CoA ligase Fat2 - Saccharomyces cerevisiae	5.9	4.1	8.4	6.8
5	Pc12g13140	strong similarity to hypothetical molasses resistency protein Rtm1 - Saccharomyces cerevisiae		1.2	1.5	1.4	0.7
	Pc12g13150	strong similarity to cdc25C associated protein kinase like protein An18g01800 - Aspergillus niger		61.4	47.6	33.1	45.3
	Pc12g13160	strong similarity to hypothetical mixed-linked glucanase precursor related protein MLG1 - Neurospora crassa		84.0	32.7	67.1	14.2
	Pc12g13170	similarity to polyketide synthase PKS17 - Botryotinia fuckeliana	Monomodular NRPS-like enzyme, 1092 AA, 3 introns, Similar enzymes in Aspergilli: A. niger (61% identity, emb CAK96299.1), oryzae (60% identity, gi 83768531 dbj BAE58668.1), A. terreus (60% identity, gi 114196109 gb EAU37809.1), A. nidulans (55% identity, gi 40745358 gb EAA64514.1) within orthologous clusters	10.1	3.6	18.8	11.4
	Pc12g13180	strong similarity to 3-dehydroshikimate dehydratase qa-4 - Neurospora crassa		15.5	19.1	18.0	21.5
	Pc12g13190	strong similarity to cDNA O-methyltransferase mt-1 - Aspergillus parasiticus		8.0	4.6	12.6	3.3
	Pc12g13200	strong similarity to hypothetical protein contig46_part_ii.tfa_3260wg - Aspergillus fumigatus		466.4	540.0	863.9	1413.8
6	Pc13g04460	strong similarity to hypothetical protein contig_1_89_scaffold_6.tfa_690wg - Aspergillus nidulans		153.2	138.1	83.2	54.4
	Pc13g04470	strong similarity to lovastatin diketide synthase lovF - Aspergillus terreus	polyketide synthase, 2591 AA, 6 introns, related enzymes in Solorina crocea (46% identity,	1.9	1.4	0.6	1.8
	Pc13g05170	similarity to multidrug resistance protein mdr1 - Aspergillus flavus		4.8	6.0	5.8	3.0
	Pc13g05180	strong similarity to dicarboxylic amino acid permease Dip5 - Saccharomyces cerevisiae		4.6	3.0	2.3	2.2
	Pc13g05190	weak similarity to 2-haloacid halido-hydrolyase IVa - Burkholderia cepacia		44.4	34.0	44.8	54.4
	Pc13g05200	strong similarity to hypothetical nuclear migration protein - Neurospora crassa		45.4	36.3	44.9	31.0
	Pc13g05210	similarity to hypothetical protein An08g09910 - Aspergillus niger		3.5	3.8	4.4	2.5
	Pc13g05220	strong similarity to microsomal cytochrome like protein An05g00300 - Aspergillus niger		729.8	864.4	786.0	1126.7
	Pc13g05230	strong similarity to ATP-dependent RNA helicase cdc28p - Schizosaccharomyces pombe		87.2	80.5	60.1	44.9
	Pc13g05240	strong similarity to protein kinase Swe1 - Saccharomyces cerevisiae		69.6	64.2	40.4	39.3
7	Pc13g05250	ferrichrome synthetase, 5081 AA, 5 introns, Similar synthetase in A. clavatus	ferrichrome synthetase, 5081 AA, 5 introns, Similar synthetase in A. clavatus (52% identity, gb EAW06285.1), which however has an intact second A-domain, thus producing presumably ferricrocin	52.7	61.2	69.2	80.5

	Pc13g05260	strong similarity to L-ornithine N5-hydroxylase psbA - Pseudomonas sp.		64.9	66.5	57.1	57.0
	Pc13g05270	strong similarity to ankyrin repeat-containing protein Akr1 - Saccharomyces cerevisiae		243.1	231.2	206.0	137.1
	Pc13g08640	strong similarity to FAD dependent L-sorbose dehydrogenase SDH - Gluconobacter oxydans		44.8	55.2	100.0	149.2
	Pc13g08650	similarity to integral membrane protein PTH11 from patent WO9913094-A2 - Magnaporthe grisea		0.2	0.6	0.6	2.2
	Pc13g08660	hypothetical protein		2.8	2.1	5.2	4.9
	Pc13g08670	weak similarity to hypothetical protein YER080w - Saccharomyces cerevisiae		4.0	2.3	4.9	2.4
	Pc13g08680	similarity to trichothecene 3-O-acetyltransferase TRI101 - Gibberella zeae		1.6	1.5	8.7	0.9
8	Pc13g08690	strong similarity to lovastatin diketide synthase lovF - Aspergillus terreus	polyketide synthase, 2526 AA, 6 introns, Related enzyme in A. oryzae (45% identity,	61.9	33.2	41.6	14.4
	Pc13g08700	weak similarity to hypothetical protein BAB11562.1 - Arabidopsis thaliana		71.0	39.9	79.6	52.0
9	Pc13g12570	strong similarity to saframycin Mx1 synthase safA - Myxococcus xanthus	Monomodular NRPS-like enzyme, 1037 AA, Similar enzymes in Aspergilli, e.g. A. oryzae (57% identity, dbj BAE64320.1), A. niger (56% identity, emb CAK38572.1)	1.0	0.2	1.6	0.6
	Pc13g12580	strong similarity to 8-amino-7-oxononanoate synthase KAPA synthase BioF - Kurthia sp.		0.4	2.5	2.4	0.9
	Pc13g14310	similarity to methyl chloride transferase AAC72357.1 - Batis maritima	methyl transferase, similar enzymes in A. niger (53% identity, emb CAK38144.1), and A. nidulans (47% identity, gb EAA58069.1),	39.4	49.1	119.3	86.8
	Pc13g14320	strong similarity to multidrug resistance protein atrD - Aspergillus nidulans [putative sequencing error]		0.7	0.2	2.4	1.5
10	Pc13g14330	weak similarity to HC-toxin peptide synthase HTS - Cochliobolus carbonum	nonribosomal peptide synthase, 5267 AA, 3 introns, Related NRPSA in A. terreus (52% identity,	3.5	2.6	7.0	6.4
	Pc14g00010	strong similarity to methionine aminopeptidase-like protein F6E13.31 - Arabidopsis thaliana		172.1	150.0	138.6	129.8
	Pc14g00020	weak similarity to hypothetical transcription regulator protein - Schizosaccharomyces pombe		85.6	132.7	153.2	343.6
	Pc14g00030	weak similarity to hypothetical protein PA1213 - Pseudomonas aeruginosa		41.5	123.1	170.5	711.3
	Pc14g00040	strong similarity to hypothetical protein nc00758.1 - Neurospora crassa		1.5	1.5	2.3	1.7
	Pc14g00050	hypothetical protein		4.7	2.9	6.0	4.2
	Pc14g00060	strong similarity to allantoin permease Dal5 - Saccharomyces cerevisiae		12.6	7.1	30.1	70.8
	Pc14g00070	hypothetical protein		1.4	2.6	4.3	3.0
11	Pc14g00080	strong similarity to nonaketide synthase lovB - Aspergillus terreus	hybrid NRPS/PKS enzyme, putative, 4049 AA, 6 introns, related NRPS/PKS in M. grisea (41 and 42%	24.4	17.5	28.6	23.0
	Pc14g00090	strong similarity to enoyl reductase of the lovastatin biosynthesis lovC - Aspergillus terreus		1.4	0.4	0.9	1.4
	Pc14g00100	similarity to hypothetical protein 162_scaffold_1.tfa_20wg - Fusarium graminearum		1.5	1.3	1.4	3.3
	Pc14g00110	weak similarity to hypothetical protein An03g00160 - Aspergillus niger		7.3	0.8	10.2	3.4
	Pc14g00120	strong similarity to cytochrome P450 trichodiene oxygenase TRI4 - Fusarium sporotrichioides		252.4	321.3	877.0	474.9
	Pc14g00130	hypothetical protein		12.9	10.1	7.9	8.8
	Pc14g00140	strong similarity to hypothetical protein contig_1_62_scaffold_4.tfa_510cg - Aspergillus nidulans		69.7	126.0	163.7	95.9
	Pc14g00150	strong similarity to lysine permease Lyp1 - Saccharomyces cerevisiae		38.7	14.5	45.6	54.0
12	Pc14g01790	strong similarity to polyketide synthase PKS17 - Botryotinia fuckeliana	Adenylate domain containing enzyme, 1075 AA, 3 introns, likely acyl substrate, Only in Ascomycetes and Basidiomycetes, related enzymes in Aspergilli, e.g. A. oryzae (50% identity, gi 83776108 dbj BAE66227.1), A. terreus (48% identity, gi 114188579 gb EAU30279.1), a related enzyme in P. nordicum (45% identity, gi 46452157 gb AAS98174.1) is involved in ochratoxin biosynthesis	1.2	3.7	2.2	1.5
	Pc15g00040	strong similarity to hypothetical protein contig17.tfa_80wg - Aspergillus fumigatus	Related to integral membrane protein in Aspergillus fumigatus (31% identity, EAL86542), Clustered with a PKS (Afu3g02570) not related to Pc15g00050	3.9	0.5	2.0	2.2
13	Pc15g00050	Polyketide synthase, 2198 AA, 2 introns, fragment, related PKS in Gibberella moniliformis	Polyketide synthase, 748 AA, 2 introns, presumably non-functional fragment, related PKS in Gibberella moniliformis (40% identity, gi 40806905 gb AAR92212.1)	1.2	0.3	0.4	3.0
	Pc15g00060	strong similarity to amino-acid N-acetyltransferase - Neurospora crassa		46.0	48.6	51.9	64.0
	Pc15g00070	similarity to arylalkylamine N-acetyltransferase AANAT1 - Esos lucius		256.8	251.5	192.8	147.1
	Pc16g00350	strong similarity to hypothetical protein An10g00160 - Aspergillus niger		1.8	2.5	5.5	1.8
	Pc16g00360	strong similarity to cytochrome P450 trichodiene oxygenase TRI4 - Fusarium sporotrichioides		0.4	1.0	1.1	3.8
14	Pc16g00370	strong similarity to 6-methylsalicylic acid synthase like protein An10g00140 - Aspergillus niger	polyketide synthase, 6-methylsalicylic acid synthase like protein, 1783 AA, also found in P. griseofulvum (85% identity, gi 1888549 gb AAB49684.1), while known 6-MSA synthases have less similarity, e.g. in Byssoscllamys nivea (48% identity, gi 13899024 gb AAK48943.1 AF360398.1), A. terreus (47% identity, gi 114213489 dbj BAA20102.2), P. griseofulvum (48% identity, gi 3212 emb CAA39295.1) related clusters in A. niger (around An10g00140), A. fumigatus (Afu8g02350), and N. fischeri (NFIA_096030).	27.5	32.1	29.4	30.2
	Pc16g00380	similarity to para-hydroxybenzoate-polyprenyltransferase ppt1p - Schizosaccharomyces pombe	Polyprenyltransferase, related enzymes in A. niger (An10g00130), N. fischeri (NFIA_096080), A. nidulans (AN8142.3), and A. fumigatus (Afu8g02350)	3.5	3.6	5.9	4.1
	Pc16g00390	similarity to hypothetical meta-cleavage compound hydrolase gene ren71 - Streptomyces aureofaciens		1.9	0.5	1.3	0.5
	Pc16g00400	strong similarity to cytochrome P450 eln2 - Coprinus cinereus		0.2	0.6	0.5	0.8
	Pc16g03740	weak similarity to siderophore biosynthesis repressor sREA - Aspergillus nidulans	High regional similarity to siderophore biosynthesis repressors sREA in Aspergilli (70 % identity)	2.2	1.8	0.6	2.2
	Pc16g03750	strong similarity to hypothetical protein contig_1_61_scaffold_4.tfa_2170cg - Aspergillus nidulans	Myosin cross reactive antigen family – various related proteins in Aspergilli (35-52% identity)	4.5	5.1	10.4	11.7

15	Pc16g03760	strong similarity to polyketide synthase PKS1 - Cochliobolus heterostrophus [putative pseudogene]	Polyketide synthase, 711 AA fragment, 50% identity to internal segments of G. zeae [gi82779926 gb ABB90283.1] and [gi42544370 gb EAA67213.1]	0.8	0.5	2.2	1.5
	Pc16g03770	strong similarity to conserved hypothetical protein - Brucella suis	Contains cupin domain, highly similar proteins in A. oryzae (88% identity, dbj BAE59540.1) and various Brucella suis sp. (88% identity, e.g. gb ABQ61818.1)	116.2	75.2	103.2	149.1
	Pc16g03780	strong similarity to enoyl reductase of the lovastatin biosynthesis lovC - Aspergillus terreus	Zn-binding dehydrogenase, related protein in A. niger (62% identity, emb CAK40659.1), Tox-D-like, e.g. in A. fumigatus (53% identity, gb EAL88863.1)	32.6	25.8	71.1	87.4
	Pc16g03790	hypothetical protein		1.0	1.3	2.8	0.3
15	Pc16g03800	strong similarity to polyketide synthase PKS1 - Cochliobolus heterostrophus	polyketide synthase, Siderophore associated, producing acyl component of triacetyl-fusarinine like siderophore, 2357 AA, 7 introns, related enzymes in A. niger (54% identity, emb CAK40660.1), and 52% identity, emb CAK41112.1))	56.8	31.4	84.8	93.7
	Pc16g03810	strong similarity to hypothetical protein F28J12.200 - Arabidopsis thaliana		1.6	2.3	5.4	6.3
	Pc16g03820	similarity to hypothetical protein 1145_scaffold_2.tfa_290cg - Fusarium graminearum		109.0	80.2	120.4	101.7
	Pc16g03830	similarity to hypothetical protein An12g06250 - Aspergillus niger		4.4	2.5	14.3	57.9
	Pc16g03840	hypothetical protein		2.0	0.6	2.4	1.9
15	Pc16g03850	similarity to peptide synthase pesA - Metarhizium anisopliae	nonribosomal siderophore peptide synthase, 1900 AA, 1 intron, highly similar synthetase in A. terreus (67% identity, gb EAU31750.1), and related NRPS in Coccioides immitis (52% identity, gb EAS36075.1) and N. crassa (50% identity, gb EAA34175.2)	17.6	11.8	11.7	25.0
	Pc16g03860	acetylase		43.5	32.4	37.5	24.8
	Pc16g03870	strong similarity to multidrug resistance protein atrD - Aspergillus nidulans		36.7	23.1	31.4	36.0
	Pc16g03880	strong similarity to hypothetical protein contig_1_7_scaffold_1.tfa_250wg - Aspergillus nidulans		61.3	78.9	45.8	75.4
	Pc16g03890	weak similarity to hypothetical protein SPAC56E4.07 - Schizosaccharomyces pombe		71.6	72.7	69.7	112.5
	Pc16g03900	strong similarity to hypothetical protein An18g01950 - Aspergillus niger [putative sequencing error]		4.1	4.6	6.1	5.5
	Pc16g03910	strong similarity to ferrioxamine B permease Sit1 - Saccharomyces cerevisiae		80.9	95.8	179.6	162.5
	Pc16g03920	weak similarity to trichodiene synthase - Gibberella pulicaris		2.0	2.8	3.3	3.7
	Pc16g04670	strong similarity to cadmium resistance protein Ycf1 - Saccharomyces cerevisiae		852.6	755.9	669.9	577.4
	Pc16g04680	hypothetical protein		6.4	5.0	7.9	7.6
16	Pc16g04690	similarity to peptide synthase pesA - Metarhizium anisopliae	nonribosomal peptide synthase, presumably cyclopentapeptide synthetase, Related NRPS systems in all Aspergilli with best regional identities between 52 and 56% (A. clavatus gb EAW07799.1), N. fischeri gb EAW22836.1), A. fumigatus gb EAL90366.1), A. oryzae dbj BAE64185.1), A. terreus gb EAL38874.1), A. niger emb CAK448225.1), A. nidulans gb EAA65335.1) and gb EAA65835.1)	597.1	563.0	394.5	408.5
	Pc16g04700	strong similarity to polyamine transport protein Tpo1 - Saccharomyces cerevisiae	Major facilitator superfamily, highly similar in Aspergilli (75-80% identity)	25.8	41.7	23.1	30.8
	Pc16g04710	strong similarity to hypothetical protein YKR018c - Saccharomyces cerevisiae	Hypothetical protein, highly similar in Aspergilli (64-69% identity)	147.0	218.2	266.9	307.8
	Pc16g04720	weak similarity to hypothetical protein YDR363w-a - Saccharomyces cerevisiae	Putative proteasome complex subunit, similar proteins in Aspergilli (65-72% identity)	201.0	240.6	238.8	265.8
	Pc16g04860	similarity to hypothetical protein An07g02510 - Aspergillus niger [putative pseudogene]	178 AA fragment, related to a hypothetical protein in A. oryzae (695 AA, regional 42% identity, dbj BAE54609.1)	1.9	1.0	0.6	2.6
	Pc16g04870	strong similarity to cytochrome P450 protein like protein An15g05070 - Aspergillus niger		0.2	0.4	4.6	5.6
	Pc16g04880	weak similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	Fungal transcription factor, similar protein in Ajellomyces capsulatus (56% identity, gb EDN11107.1), A. terreus (56% identity, gb EAU37616.1) and A. niger (54% identity, emb CAK42482.1)	375.1	389.1	381.9	377.1
17	Pc16g04890	strong similarity to polyketide synthase like protein An01g01130 - Aspergillus niger	polyketide synthase, 2546 AA, 3 introns, highly similar PKS in an orthologous cluster A. niger (66% identity, emb CAK42483.1), and PKS with related N-terminally regions (52% identity) in A. terreus (gi 114192117 gb EAU33817.1) and N. crassa (gi 28919501 gb EAA28933.1)	39.9	27.7	56.6	43.5
	Pc16g04900	strong similarity to hypothetical protein contig1477_1.tfa_1070cg - Aspergillus fumigatus	Related proteins with weak similarity in Aspergilli (31%), e.g. N. fischeri (35%, gb EAW21131.1), and 30%, gb EAW20557.1), A. fumigatus (31%, gb EAL84504.1), and gb EAL85603.1)	6.8	8.8	7.6	11.4
	Pc16g04910	strong similarity to protein involved in cercosporin production CFP - Cercospora kikuchii	Major Facilitator Superfamily, highly similar proteins in A. niger (81% identity, emb CAK42480.1) and A. terreus (80% identity, gb EAU37617.1)	5.6	18.3	12.4	7.5
18	Pc16g09930	strong similarity to FK520 biosynthetic gene cluster polyketide synthase like protein An12g02750 - Aspergillus niger	Monomodular NRPS-like enzyme, 967 AA, 1 intron, only weakly related enzymes in A. terreus (34% identity, gb EAU37830.1) and A. niger (33% identity, emb CAK41114.1)	11.3	9.6	4.7	3.8
	Pc16g09940	strong similarity to hypothetical protein 1324_scaffold_5.tfa_120cg - Fusarium graminearum	2-oxoglutarate Fe(II) oxygenase superfamily, related proteins in Sclerotinia sclerotiorum (40% identity, EDN92102) and A. clavatus (39% identity, EAW14281)	21.3	9.9	11.8	8.0
	Pc16g09950	similarity to conserved hypothetical protein - Pseudomonas syringae	Similarity to conserved hypothetical proteins in Sclerotinia sclerotiorum (41% identity, EDN92101), A. clavatus (40%, EAW14280) and Phaeosphaeria nodorum (38%, EAT86602)	5.7	5.7	3.3	3.1
	Pc16g11440	hypothetical protein	hypothetical protein, partially related proteins in Aspergilli (e.g. in A. oryzae, BAE66300.1) and A. fumigatus (EAE90389), related to superoxide dismutase	27.2	28.1	75.4	36.9
	Pc16g11450	strong similarity to hypothetical protease Mch5 - Saccharomyces cerevisiae	Monocarboxylate permease homolog, similar A. niger (CAK43292, 62% identity), A. terreus (EAL36164, 55% identity) and A. nidulans (EAA64469, 512% identity)	3.0	2.6	18.7	23.6
	Pc16g11460	strong similarity to high-affinity nicotinic acid permease Tna1 - Saccharomyces cerevisiae		128.3	89.8	142.2	351.1
	Pc16g11470	strong similarity to ATP-binding cassette multidrug transport protein atrB - Aspergillus nidulans		73.8	65.5	249.8	116.6
19	Pc16g11480	strong similarity to lovastatin diketide synthase lovF - Aspergillus terreus	polyketide synthase, 2624 AA, 7 introns, Related enzymes in N. fischeri (46% identity, gi 119406936 gb EAW16886.1), A. clavatus (46% identity, gi 119401914 gb EAW12336.1), A. oryzae (46% identity, gi 83764427 dbj BAE54571.1), A. fumigatus (46% identity, gi 66846091 gb EAL86424.1), A. terreus (41% identity, gi 66846091 gb EAL86424.1) in related clusters, presumably ochratoxin-type biosynthesis	7.3	2.7	688.5	208.1
	Pc16g11490	weak similarity to hypothetical dihydrofolate reductase CAB16576.1 - Schizosaccharomyces pombe	DUF341 domain protein, hypothetical dihydrofolate reductase, homologs in all Aspergilli, 66-71% identity	0.7	1.6	252.2	94.9
	Pc16g11500	strong similarity to hypothetical integral membrane protein - Schizosaccharomyces pombe	DUF1212 domain membrane protein, homologs in Aspergilli (60-68% identity)	82.1	79.1	644.7	345.2

20	Pc16g13880	strong similarity to hypothetical oxidoreductase PC14 - <i>Phytophthora capsici</i> [truncated ORF][putative pseudogene]	strong similarity (83% identity) to hypothetical oxidoreductase PC14 - <i>Phytophthora capsici</i> induced by interaction with host plant	2.1	1.5	1.4	3.1
	Pc16g13890	strong similarity to hypothetical membrane transport protein SPAC3H1.06c - <i>Schizosaccharomyces pombe</i>	Major facilitator superfamily, efflux pump, similar genes (53% identity) in <i>A. niger</i> (An07g00060 and An02g08330) and <i>B. fucelliana</i> (BC1G_00899)	10.2	8.1	17.4	30.4
	Pc16g13900	strong similarity to hypothetical protein Z - <i>Streptomyces hygroscopicus</i>	Putative hydrolase, related genes in Aspergilli (51-52% identity): <i>A. fumigatus</i> (EAL85114), <i>A. terreus</i> (EAL78973), <i>A. niger</i> (An07g00020), <i>A. clavatus</i> (EAW15063)	19.4	10.0	7.1	6.1
	Pc16g13910	similarity to hypothetical protein contig14.tfa_150wg - <i>Aspergillus fumigatus</i>	Putative integral membrane protein, weak similarity (34% identity) to <i>N. fischeri</i> EAW19777 and EAW 15333, <i>A. clavatus</i> (31% identity)	0.4	1.3	5.0	4.5
	Pc16g13920	strong similarity to enoyl reductase of the lovastatin biosynthesis lovC - <i>Aspergillus terreus</i>	Putative Zn-binding oxidoreductase, related enzymes in <i>N. fischeri</i> (43% identity, EAW16807; 37% identity, EAW16241), <i>A. nidulans</i> (41% identity, AN8409), <i>A. clavatus</i> (38% identity, EAW06525), <i>A. fumigatus</i> (37% identity, EAL88863)	2.1	1.0	5.6	2.4
	Pc16g13930	strong similarity to lovastatin nonaketide synthase lovB - <i>Aspergillus terreus</i>	hybrid NRPS/PKS enzyme, 4096 AA, 4 introns, related multienzymes in Aspergilli (38% identity): <i>A. clavatus</i> gij119404686[gb][EAW15062.1], <i>A. fumigatus</i> gij66844776[gb][EAL85113.1], <i>A. terreus</i> gij114197271[gb][EAL38971.1] in related clusters	103.2	88.5	68.0	52.2
	Pc16g13940	strong similarity to hypothetical protein mg09154.1 - <i>Magnaporthe grisea</i>	similarity to hypothetical proteins in <i>Magnaporthe grisea</i> (43% identity, EAQ71548), <i>Phaeosphaeria nodorum</i> (41% identity, EAT49431), <i>A. niger</i> (38% identity, CAK46265)	69.0	68.4	42.8	43.8
21	Pc18g00330	strong similarity to hypothetical phd finger transcription regulator - <i>Schizosaccharomyces pombe</i>	hypothetical phd finger transcription regulator, related proteins in <i>A. clavatus</i> (57% identity, EAW09774), <i>N. fischeri</i> (56% identity, EAW21364), <i>A. fumigatus</i> (55% identity, EAL92380)	36.0	29.3	27.4	27.3
	Pc18g00340	similarity to hypothetical protein BAB75479.1 - <i>Nostoc</i> sp.	hypothetical protein containing tetratricopeptid-domain, related proteins in <i>C. globosum</i> (46-49% identity, e.g. EAQ83821, 84130, 88760, 93491)	29.6	18.9	28.3	28.8
	Pc18g00350	hypothetical protein	hypothetical protein, weakly related proteins in <i>G. zeae</i> (28% identity, FG08961), <i>C. globosum</i> (27% identity, EAQ93177), <i>A. fumigatus</i> (28% identity, EAL85340)	108.1	80.4	140.1	91.9
	Pc18g00360	strong similarity to hypothetical protein contig336.tfa_290wg - <i>Aspergillus fumigatus</i> [putative sequencing error]	sigma-70 region 2 family protein, related proteins in all Aspergilli, e.g. <i>A. clavatus</i> (44% identity, EAW09773), <i>N. fischeri</i> (45% identity, EAW21364), <i>A. fumigatus</i> (55% identity, EAL92380)	2.8	4.0	27.5	26.4
	Pc18g00370	hypothetical protein	hypothetical protein, no significant similarity found	2.7	2.0	6.4	2.4
	Pc18g00380	strong similarity to saframycin Mx1 synthase safA - <i>Myxococcus xanthus</i>	Monomodular NRPS-like enzyme, 1318 AA, Highly similar enzymes in Aspergilli, presumably primary metabolism, <i>A. terreus</i> (75% identity, gij114195665[gb][EAU37365.1]), <i>N. fischeri</i> (73% identity, gij119407835[gb][EAW17784.1]), <i>A. fumigatus</i> (72% identity, gij66851266[gb][EAL91592.1]), <i>A. nidulans</i> (72% identity, gij66851266[gb][EAL91592.1])	393.4	398.1	386.6	366.8
	Pc18g00390	strong similarity to high-affinity glucose transporter HGT1 - <i>Kluyveromyces lactis</i>	high-affinity glucose transporter, related proteins in all Aspergilli, e.g. <i>A. clavatus</i> (57% identity, EAW09772), <i>N. fischeri</i> (65% identity, EAW21366), <i>A. fumigatus</i> (66% identity, EAL92382)	45.5	36.5	54.4	46.3
	Pc18g00400	similarity to hypothetical transcription regulator SPBC530.05 - <i>Schizosaccharomyces pombe</i>	Transcription factor, GAL4-like Zn2Cys6 binuclear cluster DNA-binding domain, related proteins in all Aspergilli, e.g. <i>A. clavatus</i> (64% identity, EAW09771), <i>N. fischeri</i> (65% identity, EAW21367), <i>A. fumigatus</i> (65% identity, EAL92384)	197.6	174.3	178.0	122.0
	Pc18g00410	strong similarity to prolyl-4-hydroxylase-alpha like protein An02g07870 - <i>Aspergillus niger</i>	prolyl-4-hydroxylase-alpha like protein, related proteins in all Aspergilli, e.g. <i>A. clavatus</i> (63% identity, EAW09770), <i>N. fischeri</i> (58% identity, EAW21368), <i>A. fumigatus</i> (62% identity, EAL92385)	8.7	8.0	10.3	11.4
	Pc18g00420	transcription factor pacC - <i>Penicillium chrysogenum</i>	transcription factor pacC, related proteins in all Aspergilli, e.g. <i>A. clavatus</i> (64% identity, EAW09769), <i>A. niger</i> (69% identity, CAK37752), <i>A. parasiticus</i> (70% identity, AF408430)	211.5	171.5	219.2	151.2
22	Pc20g02260	similarity to aminoadipate reductase enzyme lys2 - <i>Acremonium chrysogenum</i>	Monomodular NRPS-like enzyme, 1080 AA, 1 intron, Weakly related enzymes in <i>G. zeae</i> (46% identity, ref XP_382427.1), <i>Botryotinia fuckeliana</i> (45% identity, gb EDN18359.1) and <i>A. nidulans</i> (43% identity, gb EAA60062.1)	3.0	1.8	2.9	2.8
23	Pc20g02590	strong similarity to saframycin Mx1 synthase safA - <i>Myxococcus xanthus</i>	Monomodular NRPS-like enzyme, 1045 AA, related enzymes in <i>N. fischeri</i> (46% identity, gij119407216[gb][EAW17166.1]), <i>A. terreus</i> (43% identity, gij114192344[gb][EAU34044.1]), <i>A. oryzae</i> (41% identity, gij83774195[dbj][BAE64320.1])	0.5	0.9	2.1	0.8
24	Pc20g04890	similarity to salicylate hydroxylase nahG - <i>Pseudomonas putida</i>	Monoxygenase, highly similar enzymes in <i>A. oryzae</i> (80% identity, BAE64963), <i>A. terreus</i> (79% identity, EAU33487), <i>N. fischeri</i> (78% identity, EAW23864)	7.6	7.5	20.7	15.9
	Pc20g04900	strong similarity to hypothetical protein An08g08230 - <i>Aspergillus niger</i>	3123 AA, 6 introns, similar protein in <i>A. niger</i> (42% identity, emb CAK45810.1), related proteins in Aspergilli with 28-29% identity	2.7	2.2	4.2	2.2
	Pc20g04910	hypothetical protein		1.1	3.1	7.0	3.0
	Pc20g04920	similarity to fluconazole resistance protein like protein An16g02330 - <i>Aspergillus niger</i>		3.9	1.3	5.4	5.7
	Pc20g04930	similarity to polyketide synthase like protein An12g07070 - <i>Aspergillus niger</i>		0.6	0.6	3.0	1.1
	Pc20g09620	strong similarity to multidrug resistance protein Hoi1 - <i>Saccharomyces cerevisiae</i>	Major facilitator superfamily protein, related transporters in <i>Pichia guilliermondii</i> (44% identity, EDK41085), <i>Phaeosphaeria nodorum</i> (39% identity, EEEAT81452), <i>A. terreus</i> (38% identity, EAU38738)	0.8	1.5	2.7	1.1
	Pc20g09630	similarity to lipase LipP - <i>Pseudomonas</i> sp.	Putative hydrolase, lipase, similar proteins in <i>A. terreus</i> (59% identity, EAU29496), <i>A. clavatus</i> (56% identity, EAW09316), <i>N. fischeri</i> (55% identity, EAW21829)	17.6	27.9	34.8	45.4
	Pc20g09640	strong similarity to hypothetical sugar transporter encoded by DRA0271 - <i>Deinococcus radiodurans</i>	5S rRNA	139.4	129.9	134.6	113.1
	Pc20g09650	hypothetical protein	Sugar transporter, similar proteins in <i>A. terreus</i> (81% identity, EAU32826 and 66% identity, EAW29729), <i>A. oryzae</i> (71% identity, BAE60291), <i>A. clavatus</i> (65% identity, EAW21829)	0.4	0.7	1.5	1.0
	Pc20g09660	similarity to hypothetical protein required for biosynthesis of the host-specific AK-toxin Akt2 - <i>Alternaria alternata</i>		0.6	1.9	1.5	3.6
	Pc20g09670	strong similarity to hypothetical conserved protein CC0299 - <i>Caulobacter crescentus</i> [putative sequencing error]	Small noncoding RNA	0.4	0.3	2.5	0.4
	Pc20g09680	strong similarity to astaxanthin synthetase like protein An07g00250 - <i>Aspergillus niger</i>	Putative AK-toxin biosynthesis protein, related proteins in <i>N. fischeri</i> (57% identity, EAW24908), <i>A.</i>	2.3	2.0	3.1	0.6

25	Pc20g09690	strong similarity to polyketide synthase PKS17 - Botryotinia fuckeliana	Predicted phosphotransferase, related proteins in <i>A. oryzae</i> (55% identity, BAE55799), <i>N. fischeri</i> (55% identity, EAW23466), <i>A. fumigatus</i> (55% identity, EAL88000), <i>A. nidulans</i> (53% identity, AN2272), <i>A. terreus</i> (51% identity, EAU37482), <i>A. clavatus</i> (54% identity, EAW08403), <i>A. niger</i> (55% identity, An01g05030)	2.3	2.0	6.3	4.3
	Pc20g09700	hypothetical protein	P450 oxygenase, similarity to <i>A. niger</i> (44% identity, An07g00250), <i>A. terreus</i> (42% identity, EAU30769),	0.5	0.4	0.4	1.8
	Pc20g09710	strong similarity to hypothetical protein An15g06870 - Aspergillus niger	Monomodular NRPS-like enzyme, 1072 AA, 3 introns, Related enzymes in Aspergilli, e.g. <i>A. terreus</i>	166.6	111.5	48.2	51.1
	Pc20g09720	similarity to hypothetical protein An01g07110 - Aspergillus niger [putative pseudogene]	Small noncoding RNA	58.9	46.5	24.5	17.4
	Pc20g09730	similarity to glutamyl endoprotease polypeptide like protein An11g07120 - Aspergillus niger		56.2	47.0	10.8	26.6
	Pc20g09740	similarity to hypothetical zinc-finger protein SPBC15D4.02 - Schizosaccharomyces pombe	Putative phosphotransferase, related proteins in <i>Sclerotinia sclerotiorum</i> (43% identity, EDO00056)	104.3	159.8	88.6	141.2
26	Pc20g12670	strong similarity to hypothetical protein 1457_scaffold_8.tfa_130cg - Fusarium graminearum	Monomodular NRPS-like enzyme, 1283 AA, 1 intron, related enzyme in <i>A. terreus</i> (40% identity, gj114190456[gb][EAU32156.1])	3.6	0.7	5.7	5.8
	Pc21g00890	strong similarity to allantoinase permease Dnf5 - Saccharomyces cerevisiae	Major facilitator superfamily, related genes in <i>A. niger</i> (65% identity, An11g08880), <i>C. immitis</i> (61% identity, EAS28823), <i>A. terreus</i> (44% identity, EAU30298),	11.5	9.0	11.8	11.3
27	Pc21g00900	strong similarity to methicillin resistance gene HmrA - Staphylococcus aureus	Predicted peptidase, related enzymes in <i>A. oryzae</i> (67% identity, BAE55394), <i>C. immitis</i> (61% identity, EAS28824), <i>A. niger</i> (56% identity, An11g08890),	4.9	5.1	4.8	13.2
	Pc21g00910	strong similarity to acetamidase amds - Aspergillus oryzae	Predicted amidase, related enzymes in <i>A. oryzae</i> (70% identity, BAE55395), <i>A. terreus</i> (66% identity, EAU32722), <i>A. clavatus</i> (65% identity, EAW09194),	86.0	144.3	129.7	237.0
	Pc21g00920	similarity to transcription activator of lysine pathway Lys14 - Saccharomyces cerevisiae	predicted transcription activator, weakly similar proteins in <i>A. niger</i> (34% identity, An11g08910 and An03g01500), <i>A. oryzae</i> (39% identity, BAE55396)	104.9	102.1	114.0	128.9
	Pc21g00930	similarity to SR protein kinase Sky1 - Saccharomyces cerevisiae	S/T-kinase, related proteins in <i>A. clavatus</i> (77% identity, EAW11545), <i>N. fischeri</i> (76% identity, EAW16825), <i>A. terreus</i> (73% identity, EAU34098),	17.6	11.1	14.4	28.4
	Pc21g00940	strong similarity to hypothetical protein An08g05350 - Aspergillus niger	Predicted transferase, LysR family regulatory protein, related proteins in <i>N. fischeri</i> (39% identity, EAW20697 and EAW20091), <i>A. clavatus</i> (39% identity, EAW13628)	15.8	3.1	9.8	11.2
	Pc21g00950	similarity to hypothetical protein contig_1_168_scaffold_14.tfa_260wg - Aspergillus nidulans	hypothetical protein, weak similarity to <i>A. oryzae</i> (30% identity, BAE64032) and <i>A. nidulans</i> (31% identity, AN9006),	700.0	148.6	436.1	328.0
	Pc21g00960	strong similarity to lovastatin diketide synthase lovF - Aspergillus terreus	polyketide synthase, 2988 AA, 5 introns, Similar enzymes in some Aspergilli: <i>N. fischeri</i> (53% identity, gj119406436[gb][EAW16387.1]); <i>A. oryzae</i> (52% identity, gj83764443[dbj][BAE54587.1]), and 41% identity, gj83773906[dbj][BAE64031.1]), <i>A. terreus</i> (47% identity, gj114188537[gb][EAU30237.1])	81.0	16.6	138.1	116.3
	Pc21g00970	strong similarity to hex1 - Aspergillus nidulans	Woronin body protein HexA, related proteins in <i>N. fischeri</i> (60% identity, EAW17907), <i>A. terreus</i> (62% identity, EAU62125), <i>A. fumigatus</i> (53% identity, EAL91716),	2685.5	2952.6	2926.5	2448.5
	Pc21g00980	hypothetical protein		0.5	0.5	1.0	0.9
	Pc21g00990	hypothetical protein	Predicted membrane protein, similar protein in <i>Phaeosphaeria nodorum</i> (53% identity, EAT82522)	19.3	17.0	32.4	58.8
28	Pc21g01000	weak similarity to hypothetical protein SPBC1685.08 - Schizosaccharomyces pombe	Predicted protein containing PHD Zn finger, related proteins in <i>A. terreus</i> 66% identity, EAU33124), <i>A.</i>	25.9	24.2	18.8	14.9
	Pc21g01690	strong similarity to hypothetical membrane protein YOL119c - Saccharomyces cerevisiae	Putative MFS monocarboxylate transporter, related proteins in <i>A. clavatus</i> (74% identity, EAW12156), <i>A. fumigatus</i> (73% identity, EAL86614), <i>N. fischeri</i> (72% identity, EAW17188)	2.8	2.7	10.9	2.1
	Pc21g01700	similarity to mitochondrial respiratory function protein Mrf1 - Saccharomyces cerevisiae	mitochondrial Zn-dependent oxidoreductase, related proteins in <i>A. clavatus</i> (65% identity, EAW12157), <i>A. fumigatus</i> (65% identity, EAL86615), <i>N. fischeri</i> (65% identity, EAW17189)	2.8	2.7	10.9	2.1
	Pc21g01710	strong similarity to peptide synthase like protein An04g06260 - Aspergillus niger	nonribosomal peptide synthase, cyclodipeptide synthetase, 2174 AA, 1 intron, Related NRPS in <i>A.</i>	47.9	27.7	46.0	11.9
	Pc21g01720	similarity to hypothetical neutral amino acid permease - Neurospora crassa		9.0	12.6	27.1	140.3
	Pc21g01730	strong similarity to quinate transport protein qutD - Aspergillus nidulans	transmembrane neutral amino acid permease, related proteins in <i>Sclerotinia sclerotiorum</i> (44% identity, EDN93972, EDN92524 and EDN03493) and <i>Botryotinia fuckeliana</i> (42% identity, EDN48995)	49.8	36.5	73.7	75.1
29	Pc21g03820	similarity to lactone-specific esterase estf1 - Pseudomonas fluorescens [putative pseudogene]		0.9	0.6	0.5	1.8
	Pc21g03830	similarity to lactone-specific esterase estf1 - Pseudomonas fluorescens [putative pseudogene]		0.3	1.7	3.2	0.7
	Pc21g03840	similarity to hypothetical sterigmatocystin biosynthesis monooxygenase stcW - Aspergillus nidulans [putative pseudogene]		2.1	2.1	2.4	1.4
	Pc21g03850	hypothetical protein		0.6	0.1	0.5	0.4
	Pc21g03860	strong similarity to hypothetical protein An05g02110 - Aspergillus niger [putative pseudogene]		1.9	2.6	2.0	1.8
	Pc21g03870	similarity to ankyrin 3 (splice form 3) - Mus musculus		34.0	24.9	95.6	46.2
	Pc21g03880	strong similarity to hypothetical protein An05g02110 - Aspergillus niger [putative pseudogene]		0.9	0.9	2.3	2.5
	Pc21g03890	hypothetical protein		2.6	1.7	4.3	6.1
	Pc21g03900	similarity to hypothetical mixed-linked glucanase precursor related protein MLG1 - Neurospora crassa [putative pseudogene]		2.7	2.1	7.0	4.3
	Pc21g03910	similarity to hypothetical protein Smb20606 - Sinorhizobium meliloti		2.9	1.9	7.5	3.3
	Pc21g03920	strong similarity to hypothetical short chain dehydrogenase SPCC736.13 - Schizosaccharomyces pombe		0.9	1.2	6.1	12.3
	Pc21g03930	strong similarity to lovastatin diketide synthase lovF - Aspergillus terreus	polyketide synthase, related to compactin synthase, 2642 AA, 5 introns, Related enzymes in <i>A. clavatus</i>	7.4	2.9	14.5	10.4
	Pc21g03940	strong similarity to transposase Tan1 - Aspergillus niger [putative pseudogene]		2.9	1.5	4.4	5.0
	Pc21g03950	similarity to hypothetical transcription regulator SPBC530.05 - Schizosaccharomyces pombe		21.6	14.9	23.5	16.0
30	Pc21g03960	strong similarity to hypothetical oxidoreductase DR2595 - Deinococcus radiodurans	Oxidoreductase, orthologs in <i>N. fischeri</i> (58% identity, EAW19764), <i>A. terreus</i> (57% identity, EAU38660), <i>A. clavatus</i> (58% identity, EAW07338), <i>A. fumigatus</i> (55% identity, EAL85138)	1.5	2.6	4.0	5.4

30	Pc21g03970	strong similarity to alcohol dehydrogenase alkJ - <i>Pseudomonas oleovorans</i>	Predicted oxidoreductase, orthologs in <i>N. fischeri</i> (61% identity, EAW19762), <i>A. terreus</i> (56% identity, EAU36858),	4.7	1.8	5.7	6.8
	Pc21g03980	strong similarity to hypothetical protein contig_1_155_scaffold_12.tfa_40wg - <i>Aspergillus nidulans</i>	hypothetical protein, weakly related to <i>A. terreus</i> (33% identity, EAU36859), <i>A. nidulans</i> (33% identity, AN8523), <i>N. fischeri</i> (32% identity, EAW17812),	1.2	1.8	1.0	0.7
	Pc21g03990	strong similarity to polyketide synthase like protein An15g05090 - <i>Aspergillus niger</i>	polyketide synthase, 2445 AA, 5 introns, PKS in orthologous clusters in <i>Coccidioides immitis</i> (51% identity, gi 90307413 gb EAS37044.1), <i>N. fischeri</i> (39% identity, gi 119409812 gb EAW19759.1), <i>A. terreus</i> (40% identity, gi 114195156 gb EAU36856.1); weak similarity to compactin synthase in <i>P. citrinum</i> (35% identity, gi 23574645 dbj BAC20566.1)	1.7	1.2	1.0	1.9
	Pc21g04000	hypothetical protein		0.3	0.5	1.5	0.7
	Pc21g04010	similarity to 4-Hydroxyacetophenone monooxygenase hapE - <i>Pseudomonas fluorescens</i>	Predicted flavoprotein, related to <i>C. immitis</i> (48% identity, EAS37043), <i>C. globosum</i> (47% identity, EAQ92010), <i>N. fischeri</i> (43% identity, EAW19766), <i>A. terreus</i> (39% identity, EAU36862),	2.4	2.3	2.9	2.1
	Pc21g04020	weak similarity to trichothecene 3-O-acetyltransferase TRI101 - <i>Gibberella zeae</i>	Transferase family protein, weakly related to <i>A. clavatus</i> (34% identity, EAW12165), <i>M. grisea</i> (33% identity, EDJ97457), <i>N. fischeri</i> (31% identity, EAW19760),	2.0	0.7	2.0	1.9
	Pc21g04030	strong similarity to levodione reductase lvr - <i>Corynebacterium aquaticum</i>	Putative short-chain dehydrogenase, orthologs in in <i>N. fischeri</i> (64% identity, EAW19761), <i>A. terreus</i> (61% identity, EAU36857), <i>C. globosum</i> (47% identity, EAQ92004),	0.5	0.3	0.7	1.2
	Pc21g04040	strong similarity to monooxygenase paxM - <i>Penicillium oxalicum</i>	FAD-binding oxidoreductase, UbiH-related, orthologs in in <i>N. fischeri</i> (43% identity, EAW19765), <i>C. globosum</i> (42% identity, EAQ92008),	3.5	3.0	4.1	4.8
	Pc21g04050	similarity to hypothetical protein An08g03600 - <i>Aspergillus niger</i>	Hypothetical protein, related proteins in in <i>N. fischeri</i> (41% identity, EAW19768), <i>A. terreus</i> (37% identity, EAU36864), <i>C. globosum</i> (39% identity, EAQ92003),	5.2	1.7	3.7	5.1
31	Pc21g04750	similarity to aflatoxin biosynthesis regulatory protein aflR - <i>Aspergillus parasiticus</i>	Fungal transcription factor, related proteins in <i>A. oryzae</i> (61% identity, BAE63808), <i>A. terreus</i> (65% identity, EAU33888), <i>A. niger</i> (48% identity, An04g07240)	68.6	49.0	50.7	39.3
	Pc21g04760	strong similarity to cysteine dioxygenase CDO1 - <i>Rattus norvegicus</i>		109.5	23.1	157.7	78.2
	Pc21g04770	strong similarity to cytochrome-b5 reductase Mcr1 - <i>Saccharomyces cerevisiae</i>		4.8	3.4	8.5	11.3
	Pc21g04780	hypothetical protein		84.5	136.5	102.8	177.4
	Pc21g04790	similarity to hypothetical protein YER080w - <i>Saccharomyces cerevisiae</i>		95.7	87.8	96.2	86.7
	Pc21g04800	strong similarity to hypothetical protein An02g13770 - <i>Aspergillus niger</i>		22.8	35.4	44.7	56.6
	Pc21g04810	similarity to hypothetical protein contig42.tfa_2390wg - <i>Aspergillus fumigatus</i> [truncated ORF]		69.1	131.8	26.4	48.6
	Pc21g04820	similarity to precursor of triacylglycerol lipase like protein An13g00480 - <i>Aspergillus niger</i> [truncated ORF]		275.9	327.9	571.1	891.6
	Pc21g04830	strong similarity to hexokinase-like protein xprF - <i>Aspergillus nidulans</i> [putative sequencing error]		2669.1	2759.8	3620.1	3220.0
	Pc21g04840	strong similarity to polyketide synthase PKS1 - <i>Cochliobolus heterostrophus</i>	polyketide synthase, 2292 AA, 4 introns, highly similar to <i>A. niger</i> PKS (59% identity, emb CAK46336.1), related enzymes in <i>N. crassa</i> (43% identity, gi 28919463 gb EAA28899.1), <i>A. terreus</i> (43% identity, gi 114190751 gb EAS32451.1), <i>C. globosum</i> (42% identity, gi 88177264 gb EAS84732.1), <i>B. fuckeliana</i> (42% identity, gi 40787340 gb AAR90244.1) and <i>M. grisea</i> (41% identity, gi 39941356 ref XP_360215.1) and gi 39943482 ref XP_361278.1 ; 40% identity, gi 39945586 ref XP_362330.1)	3.4	1.4	8.6	11.1
32	Pc21g04850	weak similarity to hypothetical protein C25G4.2 - <i>Caenorhabditis elegans</i>		3.9	1.9	7.3	9.9
	Pc21g04860	hypothetical protein		11.9	12.0	9.7	7.0
	Pc21g05030	hypothetical protein		19.7	20.6	22.3	21.4
	Pc21g05040	hypothetical protein		7.3	6.3	10.5	12.1
	Pc21g05050	weak similarity to positive regulator qa-1F - <i>Neurospora crassa</i>		480.5	770.1	1169.9	973.6
	Pc21g05060	strong similarity to salicylate hydroxylase nahW - <i>Pseudomonas stutzeri</i>		19.9	150.1	182.0	68.3
	Pc21g05070	strong similarity to polyketide synthase like protein An09g01860 - <i>Aspergillus niger</i>	polyketide synthase, related to citrinin synthase, 2754 AA, 3 introns, similar PKS in <i>C. globosum</i> (57% identity, gi 88175714 gb EAS83182.1), related to citrinin synthase (<i>Monascus purpureus</i> , 41% identity, gi 52000443 dbj BAD44749.1)	71.6	69.0	65.5	29.9
	Pc21g05080	strong similarity to lovastatin diketide synthase lovF - <i>Aspergillus terreus</i>	polyketide synthase, 2667 AA, 5 introns, related NRPS in <i>A. terreus</i> (48% identity, gi 114190221 gb EAS31921.1), <i>C. immitis</i> (48% identity, gi 90304914 gb EAS34545.1), <i>A. nidulans</i> (47% identity, gi 40739419 gb EAA58609.1), 46% identity, gi 40746448 gb EAA65604.1), and 45% identity, gi 40745711 gb EAA64867.1), <i>A. oryzae</i> (47% identity, gi 83772350 dbj BAE62480.1)	26.4	123.9	93.8	38.4
	Pc21g05090	weak similarity to positive regulator of purine utilisation uaY - <i>Aspergillus nidulans</i>		117.8	181.5	128.6	128.1
	Pc21g05100	strong similarity to membrane protein Tpo2 - <i>Saccharomyces cerevisiae</i>		8.9	37.8	23.7	13.2
33	Pc21g05110	strong similarity to hypothetical protein contig_1_54_scaffold_3.tfa_480wg - <i>Aspergillus nidulans</i>		27.0	207.9	151.1	66.5
	Pc21g10750	strong similarity to hypothetical protein contig5_part_i.tfa_2940wg - <i>Aspergillus fumigatus</i>		168.5	186.7	169.9	264.9
	Pc21g10760	hypothetical protein		17.5	16.8	18.4	23.4
	Pc21g10770	strong similarity to aristolochene synthase Arr1 - <i>Penicillium roqueforti</i>		47.4	45.6	56.4	63.6
	Pc21g10780	weak similarity to hypothetical protein ncu00302.1 - <i>Neurospora crassa</i>		4.8	3.0	6.0	6.1
	Pc21g10790	strong similarity to HC-toxin peptide synthase HtS - <i>Cochliobolus carbonum</i>	cyclopeptide synthetase, 7287 AA, 4 introns, highly similar NRPS in <i>A. oryzae</i> (64% identity, gi 83772350 dbj BAE62480.1)	23.9	17.3	21.0	13.2
	Pc21g10800	strong similarity to acetylglutamate kinase N-acetyl-gamma-glutamyl-phosphate reductase precursor arg-6 - <i>Neurospora crassa</i>		2.6	2.6	4.0	2.7
	Pc21g10810	similarity to erythrocyte ankyrin like protein An04g07710 - <i>Aspergillus niger</i>		3.1	1.3	2.6	0.8
	Pc21g10820	strong similarity to branched-chain amino acid aminotransferase bcaT - <i>Lactococcus lactis</i>		0.2	0.5	1.9	0.5
	Pc21g10830	strong similarity to cytochrome P450 monooxygenase TRI11 - <i>Fusarium sporotrichioides</i>		3.2	2.7	3.3	5.2
34	Pc21g10840	strong similarity to fatty acid synthase alpha subunit fas2p - <i>Schizosaccharomyces pombe</i>		3.6	1.1	14.9	10.0
	Pc21g10850	strong similarity to cadmium resistance protein Ycf1 - <i>Saccharomyces cerevisiae</i>		4.0	5.1	11.3	8.7
	Pc21g10860	similarity to hypothetical protein YBR096w - <i>Saccharomyces cerevisiae</i>		4.1	3.0	7.3	3.2
	Pc21g10870	strong similarity to fatty-acyl-CoA synthase beta chain fas1p - <i>Schizosaccharomyces pombe</i>		202.5	106.7	281.6	140.2

	Pc21g10880	similarity to 7-aminosterol resistance protein Rta1 - <i>Saccharomyces cerevisiae</i>		347.4	175.0	872.2	572.7
	Pc21g12340	similarity to hypothetical transcription regulator SPAC139.03 - <i>Schizosaccharomyces pombe</i>		5.8	4.6	81.2	57.5
	Pc21g12350	strong similarity to hypothetical membrane protein YBR043c - <i>Saccharomyces cerevisiae</i>		0.4	0.4	2.0	2.4
	Pc21g12360	strong similarity to hypothetical transcriptional regulator CAF32162.1 - <i>Aspergillus fumigatus</i>		3.3	3.2	13.2	15.3
	Pc21g12370	similarity to multidrug resistance protein like protein An11g07820 - <i>Aspergillus niger</i>		0.9	0.7	0.9	1.8
	Pc21g12380	strong similarity to hypothetical neutral amino acid permease - <i>Neurospora crassa</i> [putative sequencing error]		33.2	74.7	56.8	133.9
	Pc21g12390	strong similarity to hypothetical protein B2J23.120 - <i>Neurospora crassa</i>		74.6	65.4	556.6	487.8
	Pc21g12400	similarity to hypothetical protein An11g02890 - <i>Aspergillus niger</i>		8.1	3.4	6.2	6.1
	Pc21g12410	hypothetical protein		28.4	34.4	9.4	9.0
	Pc21g12420	strong similarity to cercosporin transporter CFP - <i>Cercospora kikuchi</i>		49.7	40.2	27.0	33.7
	Pc21g12430	strong similarity to hypothetical protein contig_1_46_scaffold_3.tfa_100wg - <i>Aspergillus nidulans</i>		115.1	109.8	192.1	188.5
34	Pc21g12440	strong similarity to lovastatin diketide synthase lovF - <i>Aspergillus terreus</i>	polyketide synthase, 2397 AA, 1 intron, related PKS in <i>G. zeae</i> (60% identity, gb 42544370 gb EAA67213.1 and gb 82779926 gb ABB90283.1)	4.2	0.8	5.1	4.4
34	Pc21g12450	strong similarity to polyketide synthase wA - <i>Aspergillus nidulans</i>	polyketide synthase, 2098 AA, related PKS in <i>G. zeae</i> (56% identity, gb ABB90282.1 and 61% identity, only N-terminal region ref XP_382571.1), <i>M. grisea</i> (41% identity, gb EDK04302.1)	2.1	3.1	7.9	9.2
	Pc21g12460	strong similarity to histone acetyltransferase Gcn5 - <i>Saccharomyces cerevisiae</i> [putative pseudogene]		1.4	0.3	0.9	0.5
	Pc21g12470	weak similarity to heterokaryon incompatibility protein het-6 - <i>Neurospora crassa</i>		8.8	9.9	38.3	49.4
	Pc21g12480	strong similarity to rhamnogalacturonase B precursor rhgB - <i>Aspergillus aculeatus</i>		6.8	3.3	6.9	10.6
	Pc21g12490	hypothetical protein		142.2	152.5	176.5	200.4
	Pc21g12500	hypothetical protein		1.6	0.5	1.3	2.5
	Pc21g12510	similarity to esterase like protein An11g04020 - <i>Aspergillus niger</i> [putative pseudogene]		3.7	3.4	5.8	2.2
	Pc21g12520	similarity to hypothetical protein An08g01890 - <i>Aspergillus niger</i> [putative pseudogene]		3.0	1.8	2.4	3.2
	Pc21g12530	strong similarity to hypothetical nicotinate phosphoribosyltransferase Npt1 - <i>Saccharomyces cerevisiae</i> [putative sequencing error]		100.2	108.8	108.2	137.6
	Pc21g12540	strong similarity to hypothetical protein An01g11750 - <i>Aspergillus niger</i>		0.3	0.3	1.5	0.3
	Pc21g12550	hypothetical protein		0.6	0.3	1.3	1.0
	Pc21g12560	hypothetical protein		0.3	0.8	3.9	2.5
	Pc21g12570	similarity to hypothetical protein SPAC13C5.04 - <i>Schizosaccharomyces pombe</i>		51.1	52.7	1113.7	872.7
	Pc21g12580	hypothetical protein		0.7	2.4	2.0	1.1
	Pc21g12590	similarity to 6-hydroxy-D-nicotine oxidase 6-HDO - <i>Arthrobacter oxidans</i>		56.7	100.6	2145.6	1607.9
	Pc21g12600	strong similarity to hypothetical protein 1465_scaffold_9.tfa_470wg - <i>Fusarium graminearum</i>		585.3	599.2	2024.5	2102.3
	Pc21g12610	weak similarity to 7alpha-cephem-methoxylase subunit cmcJ - <i>Streptomyces lactamdurans</i>		15.2	18.5	567.4	386.5
	Pc21g12620	strong similarity to asparagine synthase asnB - <i>Bacillus subtilis</i>		55.4	64.3	1901.5	1359.0
35	Pc21g12630	similarity to peptide synthase pesA - <i>Metarhizium anisopliae</i>	nonribosomal peptide synthase, cyclodipeptide synthetase, 2382 AA, 2 introns, similar NRPS in <i>G.</i>	44.4	15.2	1486.7	792.0
	Pc21g12640	strong similarity to cutinase transcription factor beta CTF1b - <i>Fusarium solani</i>		114.8	123.5	122.5	178.5
	Pc21g12650	similarity to mitochondrial protein TOM6 - <i>Neurospora crassa</i>		730.2	740.5	627.9	537.8
	Pc21g12660	strong similarity to hypothetical conserved protein SPAC1952.06c - <i>Schizosaccharomyces pombe</i>		59.6	67.6	72.7	89.6
	Pc21g12670	hypothetical protein		2.2	0.8	3.5	1.1
	Pc21g12680	strong similarity to 138 kD subunit of DNA-dependent RNA polymerase II rpb2p - <i>Schizosaccharomyces pombe</i>		538.1	517.3	528.4	463.9
	Pc21g12690	strong similarity to hypothetical protein CAD21276.1 - <i>Neurospora crassa</i>		2594.8	2682.4	3200.6	2583.1
	Pc21g12700	strong similarity to hypothetical methyltransferase AAO34671.1 - <i>Gibberella zeae</i>		8.8	7.8	8.0	7.5
	Pc21g15110	hypothetical protein		0.4	0.4	0.4	1.1
	Pc21g15120	hypothetical protein		4.5	3.0	3.8	1.5
	Pc21g15130	strong similarity to hypothetical protein CAB91439.2 - <i>Neurospora crassa</i>		12.6	23.0	11.7	16.4
	Pc21g15140	strong similarity to 2,4-dihydroxyhept-2-ene-1,7-dioic acid aldolase hpcH - <i>Escherichia coli</i>		8.3	22.0	40.1	98.7
	Pc21g15150	strong similarity to hypothetical protein contig_1_139_scaffold_11.tfa_60cg - <i>Aspergillus nidulans</i>		0.3	1.5	3.5	1.5
36	Pc21g15160	strong similarity to lovastatin diketide synthase lovF - <i>Aspergillus terreus</i>	polyketide synthase, related to fumonisin synthase, 2447 AA, 17 introns, related PKS in <i>Aspergillus</i> (<i>A. oryzae</i> , 41% identity, dbj BAE54587.1 ; <i>N. fischerii</i> , 41% identity, gb EAW16387.1 ; <i>A. niger</i> , 42% identity, emb CAK43811.1 ; <i>A. clavatus</i> , 40% identity), and <i>G. moniliformis</i> (40% identity, gb AAD43562.2 AF155773_1)	0.9	0.7	3.3	2.0
	Pc21g15170	strong similarity to GTP-binding associated protein like protein An02g10400 - <i>Aspergillus niger</i> [putative sequencing error]		279.7	246.6	175.0	189.9
	Pc21g15420	strong similarity to cercosporin transporter CFP - <i>Cercospora kikuchi</i>	dimethylallyltryptophan synthase DmaW , 422 AA, 1 intron, Strong similarity in <i>Neosartorya fischerii</i> (67% identity, gb EAW17508.1), similar enzymes in <i>N. fischerii</i> (50% identity, gb EAW19789.1) and <i>A. fumigatus</i> (50% identity, gb EAL85105.2)	1629.0	807.3	2564.7	668.8
37	Pc21g15430	strong similarity to dimethylallyltryptophan synthase dmaW - <i>Claviceps purpurea</i>		883.9	595.2	2123.9	823.6
	Pc21g15440	weak similarity to hypothetical UbiE COQ5 family methyltransferase - <i>Caulobacter crescentus</i>		769.1	614.6	3302.2	1796.0
	Pc21g15450	strong similarity to cytochrome P450 monooxygenase P450I - <i>Gibberella fujikuroi</i>		344.1	281.2	1420.2	617.7
	Pc21g15460	strong similarity to maackiain detoxification protein MAK1 - <i>Nectria haematococca</i>		345.6	167.9	1829.6	653.9
	Pc21g15470	similarity to mitochondrial 25-hydroxyvitamin D3 24-hydroxylase cP450cc24 - <i>Gallus gallus</i>		241.7	115.5	742.6	98.6

37	Pc21g15480	strong similarity to cyclic peptide AM-toxin synthase like protein An03g00650 - <i>Aspergillus niger</i>	nonribosomal peptide synthase, cyclodipeptide synthetase, 2372 AA, 3 introns, related NRPS in N.	562.2	207.4	1488.2	162.2
	Pc21g15490	strong similarity to protein phosphatase PP2A0 B subunit gamma isoform - <i>Oryctolagus cuniculus</i>		109.7	97.1	90.7	83.7
38	Pc21g15990	similarity to brown 2 protein abr2 - <i>Aspergillus fumigatus</i> [putative pseudogene]		0.7	1.4	1.0	0.3
			polyketide synthase, conidial yellow pigment synthase PksP, 2138 AA, 4 introns, highly similar PKS in <i>Aspergilli</i> (<i>A. oryzae</i> , 70% identity, dbj BAE61567.1 ; <i>A. niger</i> , 68% identity, emb CAL00851.1 ; <i>A. nidulans</i> , 66% identity, emb CAA46695.2 ; <i>A. clavatus</i> , 66% identity, gb EAW14609.1 ; <i>N. fischerii</i> , 66% identity, gb EAW19338.1 ; <i>A. fumigatus</i> , 65% identity, gb AAC39471.1 ; and <i>A. nidulans</i> wA, 65% identity, prf 1905375A	8.0	4.7	12.7	4.6
	Pc21g16000	strong similarity to polyketide synthase PKS1 - <i>Colletotrichum lagenarium</i>		2.0	1.8	4.6	4.7
	Pc21g16010	hypothetical protein					
	Pc21g21240	similarity to retrotransposon like protein An11g08670 - <i>Aspergillus niger</i> [putative pseudogene]		0.1	0.1	0.8	0.1
	Pc21g21250	strong similarity to hypothetical protein An02g09120 - <i>Aspergillus niger</i> [putative pseudogene]		0.6	0.5	0.9	0.3
	Pc21g21260	weak similarity to OSJNBa0038P21.10 - <i>Oryza sativa</i> [putative pseudogene]		1.4	1.5	2.2	3.6
	Pc21g21270	strong similarity to hypothetical protein B9I2.20 - <i>Neurospora crassa</i>		3.0	1.9	2.6	1.7
	Pc21g21280	strong similarity to methyl sterol oxidase Erg25 - <i>Saccharomyces cerevisiae</i>		44.0	81.5	542.2	971.5
	Pc21g21290	strong similarity to hypothetical protein mg02069.1 - <i>Magnaporthe grisea</i>		119.4	955.7	3398.5	2598.6
	Pc21g21300	hypothetical protein		6.2	4.1	9.4	9.6
	Pc21g21310	hypothetical protein		0.3	0.4	0.7	0.6
	Pc21g21320	strong similarity to sequence 1 - unknown organism		3.9	15.0	24.8	65.1
	Pc21g21330	similarity to hypothetical protein An13g03520 - <i>Aspergillus niger</i> [putative pseudogene]		1.9	1.7	1.7	5.3
	Pc21g21340	similarity to hypothetical protein alr1138 - <i>Nostoc</i> sp.		12.1	51.8	9.3	23.1
	Pc21g21350	weak similarity to hypothetical conserved protein PA2776 - <i>Pseudomonas aeruginosa</i>		127.1	741.5	585.0	1037.2
	Pc21g21360	hypothetical protein		1.4	10.8	5.3	10.7
	Pc21g21370	acyl-coenzyme A:isopenicillin N acyltransferase (acyltransferase) AAT/PenDE	acyl-coenzyme A:isopenicillin N acyltransferase (acyltransferase) AAT/PenDE	1031.3	1827.0	1993.7	2714.6
	Pc21g21380	isopenicillin N synthase ips/PcbC	isopenicillin N synthase ips/PcbC	1312.0	2381.9	5107.0	5111.8
	Pc21g21390	alpha-aminoadipyl-cysteiny-l-valine synthetase pcbAB/acvA, 3790 AA, Isofunctional NRPS in	alpha-aminoadipyl-cysteiny-l-valine synthetase pcbAB/acvA, 3790 AA, Isofunctional NRPS in <i>A. oryzae</i> (79% identity, gi 83774189 dbj BAE64314.1), <i>A. nidulans</i> (66% identity, gi 40743782 gb EAA62968.1), <i>A. cephalosporium</i> (55% identity, gi 113315 sp P25464 ACVS_CEPAC), and <i>Kallichroma thetys</i> (53% identity, gi 13398458 gb AAK21902.1 AF335329_1), bacterial ACV synthetases have more than 40% identity	1056.7	2068.8	4321.8	4376.0
	Pc21g21400	hypothetical protein		0.9	1.7	2.6	2.0
	Pc21g21410	strong similarity to hypothetical protein ncu05765.1 - <i>Neurospora crassa</i>		2.6	1.7	5.4	6.3
	Pc21g21420	strong similarity to hypothetical protein An09g04870 - <i>Aspergillus niger</i>		8.8	5.6	87.6	107.0
40	Pc21g22530	strong similarity to hypothetical protein contig_1_135_scaffold_11.tfa_110wg - <i>Aspergillus</i>	Monomodular NRPS-like enzyme, 878 AA, 2 introns, unique, only distantly related domains in A.	7.8	3.2	1.3	2.5
41	Pc21g22650	strong similarity to polyketide synthase PKS17 - <i>Botryotinia fuckeliana</i>	Monomodular NRPS-like enzyme, 1225 AA, 1 intron, Highly similar enzymes in <i>Aspergilli</i> , <i>A. terreus</i>	23.4	16.9	14.3	18.2
	Pc22g08110	strong similarity to cytochrome P450 eln2 - <i>Coprinus cinereus</i>		31.6	12.9	108.9	81.6
	Pc22g08120	similarity to hypothetical meta-cleavage compound hydrolase gene ren71 - <i>Streptomyces aureofaciens</i> [putative sequencing error]		1.3	1.1	8.8	11.5
	Pc22g08130	weak similarity to protein involved in cercosporin production CFP - <i>Cercospora kikuchii</i> [putative pseudogene]		1.7	0.2	1.6	2.4
	Pc22g08140	weak similarity to hypothetical transcription regulator SPBC530.05 - <i>Schizosaccharomyces pombe</i>		121.6	106.9	86.5	90.4
	Pc22g08150	strong similarity to cytochrome p450 related protein eln2 - <i>Coprinus cinereus</i>		0.2	1.7	1.9	1.3
	Pc22g08160	strong similarity to hypothetical protein An10g00180 - <i>Aspergillus niger</i>		3.5	2.2	3.5	5.4
	Pc22g08170	strong similarity to 6-methylsalicylic acid synthase like protein An10g00140 - <i>Aspergillus niger</i>	6-methylsalicylic acid synthase, 1776 AA, 1 intron, 6-MSA synthases from <i>P. griseofulvum</i> (89% identity, emb CAA39295.1), <i>A. clavatus</i> (77% identity, gb EAW11667.1), <i>Byssoschlamys nivea</i> (73% identity, gb AAK48943.1 AF360398_1)	3.2	2.8	3.3	3.0
	Pc22g08180	strong similarity to hypothetical protein mg05770.1 - <i>Magnaporthe grisea</i>		192.5	221.2	159.4	184.7
	Pc22g08190	similarity to retroviral-like transposable element like protein An03g03110 - <i>Aspergillus niger</i> [truncated ORF]		2.1	3.0	17.5	15.1
	Pc22g08200	hypothetical protein		0.3	0.6	1.1	0.4
	Pc22g08210	weak similarity to hypothetical protein - <i>Plasmodium falciparum</i>		2.5	1.3	2.2	1.8
	Pc22g08220	strong similarity to hypothetical protein contig14.tfa_150wg - <i>Aspergillus fumigatus</i>		59.8	61.5	101.3	117.8
	Pc22g08230	hypothetical protein		1.8	1.0	8.6	2.7
	Pc22g08240	strong similarity to hypothetical protein - <i>Danio rerio</i>		12.6	8.8	44.9	50.6
	Pc22g08250	strong similarity to protein involved in cercosporin production CFP - <i>Cercospora kikuchii</i>		2.6	1.7	2.5	1.6
	Pc22g08260	similarity to aminotriazole resistance protein Atr1 - <i>Saccharomyces cerevisiae</i>		1.1	0.6	1.6	1.4
43	Pc22g09430	similarity to nonribosomal peptide synthase MxaA - <i>Stigmatella aurantiaca</i>	Monomodular NRPS-like enzyme, 1030 AA, 3 introns, Related enzymes in <i>Ajellomyces capsulatus</i> (54% identity, gb EDN05769.1), <i>A. oryzae</i> (49% identity, dbj BAE66286.1), <i>A. clavatus</i> (44% identity, gb EAW07472.1)	696.1	129.9	108.3	21.4
	Pc22g20360	strong similarity to siderophore-iron transporter for enterobactin Entb1 - <i>Saccharomyces cerevisiae</i>		1.9	4.3	19.4	67.3
	Pc22g20370	strong similarity to carnitine racemase like protein An03g03550 - <i>Aspergillus niger</i>		0.7	1.6	4.9	43.3
	Pc22g20380	strong similarity to aerobactin biosynthesis protein iucB - <i>Escherichia coli</i>		6.5	15.0	32.7	90.2

44	Pc22g20390	strong similarity to multidrug resistance protein atrD - <i>Aspergillus nidulans</i>		15.2	22.1	33.3	100.0
	Pc22g20400	weak similarity to cyclic peptide AM-toxin synthase AMT - <i>Alternaria alternata</i>	triacylfusarinin synthetase, 2076 AA, 1 intron, Highly similar NRPS in <i>N. fischerii</i> (68% identity, gb EAW17196.1), <i>A. fumigatus</i> (69% identity, gb EAL86624.1 , and 68% identity, gb AAX11421.1), <i>A. oryzae</i> (72% identity, dbj BAE65630.1 ,dbj BAE65630.1 , and 71% identity, dbj BAC78651.1), <i>A. clavatus</i> (68% identity, gb EAW12140.1), <i>A. niger</i> (65% identity, emb CAK47792.1), <i>A. nidulans</i> (62% identity, gb EAA58620.1), <i>Botryotinia fuckeliana</i> (67% identity, gb EDN31507.1 , and 65% identity, gb AAX11420.1), <i>Sclerotinia sclerotiorum</i> (68% identity, gb EDO01775.1)	16.1	18.6	34.2	77.4
	Pc22g20410	strong similarity to hypothetical protein contig43.tfa_360wg - <i>Aspergillus fumigatus</i>		29.6	46.1	73.1	151.5
	Pc22g20420	weak similarity to exo-alpha-sialdase - <i>Trypanosoma cruzi</i>		289.3	251.7	157.7	111.6
45	Pc22g22580	strong similarity to hypothetical protein contig_1_153_scaffold_12.tfa_490wg - <i>Aspergillus nidulans</i>	adenylate domain containing protein, 797 AA, 1 intron, related proteins in <i>A. nidulans</i> (47% identity, gb EAA67055.1), <i>N. fischerii</i> (45% identity, gb EAW24710.1), <i>A. terreus</i> (42% identity, gb EAU30093.1), <i>C. globosum</i> (42% identity, gb EAQ91494.1)	5.9	3.3	33.1	4.3
	Pc22g22590	similarity to alcohol dehydrogenase par2 - <i>Mus musculus</i>		17.3	71.7	54.2	44.7
	Pc22g22600	weak similarity to hypothetical protein B24P7.350 - <i>Neurospora crassa</i>		4.6	0.8	71.9	14.7
	Pc22g22610	similarity to cytochrome P450 monooxygenase avnA - <i>Aspergillus parasiticus</i>		5.9	2.2	44.6	10.9
46	Pc22g22820	strong similarity to cytochrome P450 monooxygenase P450l - <i>Gibberella fujikuroi</i>		0.6	1.0	11.8	0.7
	Pc22g22830	hypothetical protein		1.5	1.3	3.0	2.8
	Pc22g22840	strong similarity to ATP-binding cassette multidrug transport protein atrA - <i>Aspergillus nidulans</i>		2.9	1.7	3.9	4.4
	Pc22g22850	strong similarity to polyketide synthase like protein An09g01860 - <i>Aspergillus niger</i> [putative sequencing error]	polyketide synthase, 2522 AA, 4 introns, related PKS in <i>A. nidulans</i> (53% identity, gb EAA67005.1), and <i>A. terreus</i> (53% identity, gb EAU29529.1)	6.4	3.3	13.1	9.1
	Pc22g22860	similarity to cinnamyl-alcohol dehydrogenase CAD1 - <i>Eucalyptus gunnii</i>		0.7	0.6	5.1	2.1
	Pc22g22870	strong similarity to levodione reductase lvr - <i>Corynebacterium aquaticum</i>		0.5	1.5	3.4	1.0
	Pc22g22880	similarity to para-hydroxybenzoate--polyprenyltransferase ppt1p - <i>Schizosaccharomyces pombe</i>		2.0	0.6	2.2	2.3
	Pc22g22890	strong similarity to hypothetical monooxygenase paxM - <i>Penicillium paxilli</i>		0.7	0.3	2.5	2.0
47	Pc22g23700	weak similarity to fructosyl amine oxygen oxidoreductase - <i>Aspergillus fumigatus</i>		12.4	15.1	32.4	52.0
	Pc22g23710	strong similarity to pyrroline-5-carboxylate reductase P5CR - <i>Zalerion arboricola</i>		10.4	14.4	25.3	37.3
	Pc22g23720	strong similarity to hypothetical membrane transport protein SPAC3H1.06c - <i>Schizosaccharomyces pombe</i>		23.2	11.3	19.7	6.9
	Pc22g23730	hypothetical protein		2.6	0.4	1.3	2.5
	Pc22g23740	strong similarity to protein involved in betaine lipid production btaA like protein An02g02840 - <i>Aspergillus niger</i>		6.5	4.4	11.0	7.2
	Pc22g23750	strong similarity to lovastatin diketide synthase lovF - <i>Aspergillus terreus</i>	polyketide synthase, 3061 AA, 12 introns, N-terminal part has weak similarity to PKS in <i>N. crassa</i> (32% identity, gb EAA28933.1), <i>A. terreus</i> (31% identity,gb EAU33817.1), <i>A. clavatus</i> (32% identity, gb EAW08895.1), <i>G. moniliformis</i> (31% identity, gb AAD43562.2 AF155773.1)	1.8	1.6	4.3	1.4

@ Values given are the average of three independent experiments

Supplementary Table 5. ORFs on the amplified region of industrial *Penicillium* strains

orf code	Description of putative <i>P. chrysogenum</i> ORF	Best blast homolog to putative <i>P. chrysogenum</i> ORF *			Average transcript levels @				mRNA ratio #		mRNA ratio \$	
		Description (Species)	Accession number	e-value	WS - PAA	WS + PAA	DS - PAA	DS + PAA	WS	DS17690	non-producing	producing
Pc21g21240	similarity to retrotransposon like protein An11g08670 - Asperg	strong similarity to retrotransposon Tst1 of <i>Ne</i>	At11c0280	4e-62	0.1	0.1	0.8	0.1	0.8	0.1	6.3	1.0
Pc21g21250	strong similarity to hypothetical protein An02g09120 - Asperg	strong similarity to hypothetical retrotransposon	An02g0270	1e-28	0.6	0.5	0.9	0.3	0.9	0.3	1.6	0.5
Pc21g21260	weak similarity to OS-NBha008921.10 - <i>Oryza sativa</i> [putative]	gene - OS-NBha008921.10 - <i>Oryza sativa</i> [putative]	OS-NB0241_10	2e-05	1.4	1.5	2.2	1.6	1.1	1.6	1.5	2.3
Pc21g21270	strong similarity to hypothetical protein B92.20 - <i>Neurospora</i>	similarity to hypothetical protein B92.20 - <i>Neu</i>	An09g23170	3e-48	3.0	1.9	2.6	1.7	0.6	0.6	0.9	0.9
Pc21g21280	strong similarity to methyl sterol oxidase Erg25 - <i>Saccharomy</i>	probable c-4 methyl sterol oxidase - fission	T38986	1e-89	44.0	81.5	542.2	971.5	1.9	1.8	12.3	11.9
Pc21g21290	strong similarity to hypothetical protein mg02069.1 - <i>Magnapor</i>	probable laccase/guaiacolase (EC 4.4.1.8)	AG3272	1e-23	119.4	955.7	3395.5	2598.4	8.0	0.8	28.5	2.7
Pc21g21300	hypothetical protein	-	-	-	6.2	4.1	9.4	9.6	0.7	1.0	1.5	2.4
Pc21g21310	hypothetical protein	unnamed protein product [Aspergillus oryzae]	BAE61884.1	4e-10	0.3	0.4	0.7	0.6	1.3	0.8	2.1	1.3
Pc21g21320	similar to AA sequence:TREMBL-AX100446.1	Sequence 1 from Patent WO0121773 (Penicil)	AX100446_1	5e-72	3.9	15.0	24.8	65.1	3.8	2.6	6.3	4.3
Pc21g21330	similarity to hypothetical protein encoded in 13g03520 - <i>Aspergillus nig</i>	strong similarity to hypothetical protein encoded in 13g03520	-	1e-15	1.9	1.7	5.3	0.9	3.0	0.9	3.2	0.2
Pc21g21340	similarity to hypothetical protein alt1138 - <i>Neotoc sp.</i>	unnamed protein product [Aspergillus oryzae]	BAE61886.1	0.0	12.1	51.8	9.3	23.1	4.3	2.5	0.8	0.4
Pc21g21350	weak similarity to hypothetical conserved protein PAZ776 - <i>Pe</i>	similarity to hypothetical oxidoreductase PA53	An03g05870	1e-60	127.1	741.5	585.0	1037.2	5.8	1.8	4.6	1.4
Pc21g21360	hypothetical protein	-	-	-	1.4	19.5	5.3	10.7	7.6	2.0	3.7	1.0
Pc21g21370	acyl-coenzyme A isopenicillin N acyltransferase (acyltransfer	acyl-CoA6-aminopenicillanic acid acyltransfer	JQ0118	0.0	1031.3	1527.0	1993.7	2714.6	1.8	1.4	1.9	1.5
Pc21g21380	isopenicillin N synthase ipcPcbC - <i>Penicillium chrysogenum</i>	isopenicillin N synthase (EC 1.14.11.3) ipcPcb	S04441	0.0	1312.0	2381.9	5107.0	5111.8	1.8	1.0	3.9	2.1
Pc21g21390	alpha-aminoacyl-L-cysteinyl-valine synthetase pcbAB acvA -	alpha-aminoacyl-L-cysteinyl-valine synthetase	YGPLV8	0.0	1098.0	2060.0	4345.2	4557.4	1.9	1.0	4.0	2.2
Pc21g21400	hypothetical protein	-	-	-	0.9	1.7	2.5	2.0	1.9	0.8	2.8	1.2
Pc21g21410	strong similarity to hypothetical protein ncu05765.1 - <i>Neurosp</i>	unnamed protein product [Aspergillus oryzae]	BAE66036.1	5e-134	2.6	1.7	5.4	6.3	0.6	1.2	2.1	3.7
Pc21g21420	strong similarity to hypothetical protein An09g04870 - Asperg	weak similarity to polypeptide SEQ (D NO-618)	An09g04870	0.0	8.8	5.6	87.6	107.0	0.6	1.2	9.9	19.2
Pc21g21430	similarity to hypothetical protein An09g03770 - <i>Aspergillus nig</i>	hypothetical protein CHGG_05485 (Chaetom	XP_001221560.1	2e-16	4.4	3.2	22.5	41.3	0.7	1.8	5.1	12.8
Pc21g21440	hypothetical protein	-	-	-	20.0	9.1	533.7	170.9	0.5	0.3	26.7	18.8
Pc21g21450	strong similarity to hypothetical protein contig_1_94_scaffold	hypothetical protein [Aspergillus nidulans_F	contig_1_94_scaffold_6.fa	0.0	2.2	0.8	1.3	0.7	0.4	0.5	0.6	0.8
Pc21g21460	hypothetical protein	-	-	-	121.7	87.8	711.9	614.2	0.7	0.9	5.8	7.0
Pc21g21470	strong similarity to hypothetical protein contig_1_94_scaffold	hypothetical protein [Aspergillus nidulans_F	contig_1_94_scaffold_6.fa	1e-165	112.6	92.1	700.9	694.7	0.8	1.0	6.2	7.5
Pc21g21480	hypothetical protein	-	-	-	27.9	22.1	90.9	55.5	0.8	0.6	3.3	2.5
Pc21g21490	hypothetical protein	-	-	-	4.4	3.2	22.0	25.3	0.7	1.1	5.0	7.9
Pc21g21500	strong similarity to hypothetical isomyl alcohol oxidase mref	FAD binding domain protein [Neosartorya f	XP_001261742	0.0	4.9	4.3	2.7	6.1	0.9	2.3	0.5	1.4
Pc21g21510	strong similarity to ceresopitin transporter CTP - <i>Cercospor</i>	strong similarity to ceresopitin transporter CTP	At01g11290	8e-93	0.1	0.2	0.5	0.8	1.3	1.5	4.0	5.0
Pc21g21520	similarity to C-8,7 sterol isomerase - <i>Arabidopsis thaliana</i>	strong similarity to emopamil-binding protein	An01g13260	8e-56	24.9	7.9	48.6	133.8	0.3	2.8	2.0	16.9
Pc21g21530	strong similarity to allantoinase permease Dal5 - <i>Saccharomy</i>	strong similarity to allantoinase permease Dal5	An12g01960	1e-180	0.7	1.2	3.2	3.6	1.8	1.1	4.8	3.1
Pc21g21540	strong similarity to lipase Ipl1 - <i>Geotrichum candidum</i>	strong similarity to triacylglycerol lipase Ipl	An03g02270	1e-178	6.8	12.4	27.8	33.8	1.9	1.9	4.1	4.3
Pc21g21550	strong similarity to beta transducin-like protein like protein An	similarity to beta transducin-like protein heta	An01g01380	1e-163	1.4	0.8	0.2	0.8	0.6	3.3	0.2	1.0
Pc21g21560	similarity to histidine kinase like protein An07g08100 - Asperg	strong similarity to histidine kinase nik-1 of <i>Ne</i>	An07g08100	1e-11	1.5	0.9	1.3	0.7	0.6	0.6	0.8	0.8
Pc21g21570	strong similarity to hypothetical impala transposase - <i>Fusarium</i>	strong similarity to hypothetical impala transpo	An03g03020	2e-25	77.0	82.1	84.1	99.1	1.1	1.2	1.1	1.2
Pc21g21580	strong similarity to hypothetical protein encoded An04g09900 - Asperg	strong similarity to hypothetical protein encoded	An04g09900	9e-45	147.8	147.8	104.9	102.7	0.7	0.8	0.8	1.0
Pc21g21590	strong similarity to glucose transporter rco-3 - <i>Neurospora cr</i>	strong similarity to monosaccharide transport	An15g03940	1e-139	665.7	166.9	88.1	29.8	0.3	0.3	0.1	0.2
Pc21g21600	hypothetical protein	-	-	-	5.3	2.0	1.6	3.6	0.4	2.2	0.3	0.7
Pc21g21610	strong similarity to P-type ATPase ENA1 - <i>Schwanniomyces</i>	strong similarity to P-type ATPase ENA1 of <i>Sch</i>	An02g00890	0.0	-	-	-	-	-	-	-	-
Pc21g21620	weak similarity to retrotransposon Tst1 - <i>Nicotiana tabacum</i>	strong similarity to retrotransposon Tst1 of <i>Ne</i>	An10g07040	1e-106	0.1	0.3	1.5	0.5	3.0	0.4	15.0	1.8
Pc21g21630	similarity to hypothetical protein B24P11.150 - <i>Neurospora cr</i>	strong similarity to hypothetical protein B24P11.150	An16g05580	7e-16	0.7	2.3	3.9	2.3	0.4	0.6	5.8	1.0
Pc21g21640	similarity to 7-aminocholesterol resistance protein Rta1 - <i>Sac</i>	similarity to 7-aminocholesterol resistance pr	An14g07050	1e-101	1.3	0.7	3.1	3.4	0.6	1.1	2.4	4.6
Pc21g21650	strong similarity to alcohol dehydrogenase ADH like protein An	strong similarity to alcohol dehydrogenase AD	An01g14780	1e-147	6.2	6.9	7.6	6.5	1.1	0.9	1.2	1.0
Pc21g21660	weak similarity to zinc-finger transcription factor armd4 - Aspe	strong similarity to zinc-finger transcription fac	An01g14770	0.0	16.4	15.8	20.3	21.2	1.0	1.0	1.3	1.3
Pc21g21670	weak similarity to hypothetical RNA polymerase II transcriptio	similarity to gene expression regulator At14	An14g07040	8e-91	2.8	1.0	3.9	2.1	0.4	0.5	1.4	2.0
Pc21g21680	strong similarity to hypothetical C6 hypothetical protein CAD3	strong similarity to hypothetical C6 hypothetical	An09g07710	8e-52	71.9	63.9	42.1	16.3	0.9	0.4	0.6	0.3
Pc21g21690	strong similarity to ATP-dependent RNA helicase like protein	strong similarity to ATP-dependent RNA helicase	An03g07810	0.0	108.9	99.0	96.6	61.2	0.9	0.6	0.9	0.6

* Best non-*Penicillium chrysogenum* blast hits are given

@ Values given are the average of three independent experiments

Interesting ratio's are indicated in blue (i.e. ratio >2 and one of the values above the background of 12)

\$ Interesting ratio's are indicated in orange (i.e. ratio >2 and one of the values above the background of 12)

Supplementary Table 6. Putative β -lactam related ORFs located outside the amplified region of industrial *Penicillium* strains

orf code	Description of putative <i>P. chrysogenum</i> ORF	Identification method	e-value	Average transcript levels @				mRNA ratio #		mRNA ratio \$		length [aa]	Mw [Da]	pI	CAI
				WIS - PAA	WIS + PAA	DS - PAA	DS + PAA	WIS	DS17690	non-producing	producing				
Pc12g04030	strong similarity to cephalosporin esterase - Rhodospiridium toruloides	blastP with cephalosporin esterase	6e-33	3.1	2.3	3.9	1.9	0.8	0.5	1.3	0.8	653	70940	6.07	0.383
Pc12g11540	similarity to isopenicillin N epimerase celD - Streptomyces clavuligerus	blastP with Streptomyces clavuligerus celD	3.4e-16	272.6	431.6	865.3	1032.6	1.6	1.3	3.0	2.4	476	53175.8	6.14	0.477
Pc12g13400	strong similarity to cephalosporin esterase - Rhodospiridium toruloides	blastP with cephalosporin esterase	1e-40	159.5	214.3	104.2	86.3	1.3	0.7	0.7	0.3	542	59792.3	5.22	0.421
Pc13g04050	strong similarity to hypothetical protein rpgA - Aspergillus nidulans	blastP with Aspergillus nidulans rpgA	2.6e-51	144.9	117.9	139.7	160.2	0.8	1.1	1.0	1.4	422	47080.4	5.76	0.514
Pc13g04140	strong similarity to precursor of cephalosporin esterase - Rhodospiridium toruloides	blastP with cephalosporin esterase	1e-56	6.1	5.3	9.3	14.0	0.9	1.5	1.5	2.6	542	60032.6	6.15	0.442
Pc13g04180	strong similarity to β -lactamase subunit cmcJ - Streptomyces clavuligerus	blastP with Streptomyces clavuligerus cmcJ	1.5e-23	15.4	6.5	11.0	47.6	0.4	4.3	0.7	7.0	314	35521.7	6.01	0.493
Pc13g04680	strong similarity to β -lactamase subunit cmcJ - Streptomyces clavuligerus	blastP with Streptomyces clavuligerus cmcJ	5.9e-19	93.5	100.3	178.5	249.5	2.0	1.4	1.9	1.3	296	33556.6	5.6	0.443
Pc13g09140	strong similarity to isopenicillin N acyltransferase aAT - Aspergillus nidulans	blastP with Penicillium chrysogenum aAT	1e-43	1.7	2.2	2.1	1.3	1.3	0.6	1.3	0.6	362	39525.3	5.4	0.440
Pc13g12270	strong similarity to 4-coumarate-CoA ligase 4CL - Arabidopsis thaliana	blastP with Penicillium chrysogenum pclA	2.4e-103	92.2	100.8	57.3	133.4	1.1	2.3	0.6	1.3	556	61635.9	7.61	0.454
Pc13g14300	similarity to hypothetical beta-lactamase XF1621 - Xylella fastidiosa	blastP with beta-lactamase	2e-21	77.1	40.1	120.9	116.3	0.6	1.0	1.6	2.4	570	63924.9	5.8	0.407
Pc15g00420	strong similarity to very long-chain fatty acyl-CoA synthase Fat1	blastP with Acromonium chrysogenum celD1	2e-77	3.1	1.5	3.6	8.4	0.5	2.3	1.2	5.7	634	69269.2	7.56	0.508
Pc16g14010	strong similarity to hypothetical methyltransferase AAO34671.1 - Gibberella zeae	blastP with Aspergillus nidulans laeA	2.4e-145	365.6	361.9	373.8	395.3	1.0	1.1	1.0	1.1	427	48548.5	6.98	0.460
Pc16g14410	strong similarity to β -lactamase subunit cmcJ - Streptomyces clavuligerus	blastP with Streptomyces clavuligerus cmcJ	2.5e-16	13.0	13.9	12.2	16.8	1.1	1.4	0.9	1.2	262	32241.8	6.8	0.408
Pc18g05710	strong similarity to very long-chain fatty acyl-CoA synthase Fat1	blastP with Acromonium chrysogenum celD1	1e-100	79.2	84.9	103.7	101.3	1.1	1.0	1.3	1.2	658	73218.7	8.61	0.523
Pc20g04080	strong similarity to gene involved in the regulation of penicillin biosynthesis suAprgA1 - Aspergillus nidulans	blastP with Aspergillus nidulans suAprgA1	2e-82	545.4	568.9	444.0	394.5	1.0	0.9	0.8	0.7	285	32198.9	4.46	0.773
Pc20g10590	similarity to hypothetical beta-lactamase XF1621 - Xylella fastidiosa	blastP with beta-lactamase	1e-26	75.8	37.5	48.8	29.0	0.5	0.6	0.6	0.8	560	61921.3	6.58	0.390
Pc20g13500	strong similarity to very long-chain fatty acyl-CoA synthase Fat1	blastP with Acromonium chrysogenum celD1	1e-171	54.7	25.9	45.7	143.7	0.5	3.1	0.8	6.5	632	70529.4	7.01	0.478
Pc21g12610	weak similarity to β -lactamase subunit cmcJ - Streptomyces clavuligerus	blastP with Streptomyces clavuligerus cmcJ	0.019	15.2	18.5	567.4	386.5	1.2	0.7	37.4	20.9	392	43818.3	5.53	0.399
Pc21g18210	strong similarity to homoserine O-acetyltransferase metE - Aspergillus nidulans	blastP with Acromonium chrysogenum celG	7e-88	179.8	176.4	224.8	209.0	1.0	0.9	1.3	1.2	499	54534.8	5.27	0.565
Pc21g20500	strong similarity to 4-coumarate-CoA ligase 4CL - Arabidopsis thaliana	blastP with Penicillium chrysogenum pclA	2.8e-86	72.6	62.9	52.8	123.1	0.9	2.3	0.7	2.0	566	62476.9	6.57	0.463
Pc21g22010	strong similarity to 4-coumarate-CoA ligase 4CL - Arabidopsis thaliana	blastP with Penicillium chrysogenum pclA	2.1e-69	50.8	45.3	22.0	11.7	0.9	0.5	0.4	0.3	560	61901.3	6.32	0.417
Pc22g00550	similarity to hypothetical beta-lactamase XF1621 - Xylella fastidiosa	blastP with beta-lactamase	2e-35	20.1	27.6	23.1	36.2	1.4	1.6	1.1	1.3	511	56664.7	6.5	0.432
Pc22g13680	strong similarity to hypothetical protein contig1495_1 fta_1650cg - Aspergillus fumigatus	blastP with Acromonium chrysogenum celD2	1e-143	17.3	13.7	12.3	15.4	0.8	1.3	0.7	1.1	377	41017	5.86	0.431
Pc22g14900	phenylacetyl-CoA ligase pclA - Penicillium chrysogenum	blastP with Penicillium chrysogenum pclA	0.0	108.5	140.0	222.5	381.6	1.3	1.7	2.1	2.7	578	62629.2	8.63	0.489
Pc22g16570	strong similarity to serine O-acetyltransferase cysA - Aspergillus nidulans	blastP with Acromonium chrysogenum celG	2e-39	196.2	242.1	280.8	422.6	1.2	1.5	1.4	1.7	529	57623.2	6.53	0.493
Pc22g20270	strong similarity to 4-coumarate-CoA ligase 4CL - Arabidopsis thaliana	blastP with Penicillium chrysogenum pclA	2.7e-103	111.8	173.0	186.3	458.8	1.6	2.5	1.7	2.6	562	62639.9	8.46	0.537
Pc22g24630	similarity to protein involved in cephalosporin C biosynthesis like protein An09g06420 - Aspergillus nidulans	blastP with Acromonium chrysogenum celD2	2.4e-22	3.8	5.4	6.0	7.0	1.4	1.2	1.6	1.3	118	12524	5.41	0.407
Pc22g24780	strong similarity to 4-coumarate-CoA ligase 4CL - Arabidopsis thaliana	blastP with Penicillium chrysogenum pclA	3.9e-80	11.6	44.1	18.2	250.3	3.8	13.9	1.6	5.7	549	60915.1	7.2	0.436

@ Acromonium chrysogenum celD2 gene for isopenicillin N-CoA epimerase
@ Values given are the average of three independent experiments
Interesting ratio's are indicated in blue (i.e. ratio >2 and one of the values above the background of 12)
\$ Interesting ratio's are indicated in orange (i.e. ratio >2 and one of the values above the background of 12)

Supplementary Table 7. *In silico* identified putative *P. chrysogenum* microbody matrix proteins

ORF code	Description of putative <i>P. chrysogenum</i> ORF	Average transcript levels				Putative	Putative
		WIS - PAA	WIS + PAA	DS - PAA	DS + PAA	PTS1	PTS2
Pc04g00160	hypothetical protein	4.87	5.23	9.13	4.53		KLRCISCHL
Pc04g00190	hypothetical protein	9.37	6.60	15.73	9.95	CKI	
Pc06g01180	strong similarity to probable acyl-CoA dehydrogenase - Glomus intraradices	0.93	0.53	4.07	0.48	AKI	
Pc06g02280	strong similarity to cadmium resistance protein - Saccharomyces cerevisiae	435.60	379.10	427.67	409.90	SKI	
Pc07g00070	hypothetical protein	na	na	na	na	CKI	
Pc09g00320	hypothetical protein	na	na	na	na	CKI	
Pc12g00910	strong similarity to hypothetical protein AACU01001693_2 - Magnaporthe grisea	19.87	24.37	38.83	56.23		RLSRLSKHL
Pc12g01170	strong similarity to hypothetical protein Afu2g14850 - Aspergillus fumigatus	145.40	92.33	183.83	187.33	AKL	
Pc12g01670	similarity to ankyrin repeat protein E3_19 - Synthetic construct	4.67	2.73	4.33	1.80	AKM	
Pc12g02020	similarity to delta-6 desaturase like protein An07g06770 - Aspergillus niger	0.40	1.70	2.47	1.88		RLGVVAGQL
Pc12g04590	similarity to beta transducin-like protein het-e1 - Podospora anserina	482.53	499.20	514.20	486.03		RLTELLDQL
Pc12g05400	strong similarity to citrate synthase citA - Aspergillus niger	665.60	586.47	417.17	266.80	AKL	
Pc12g05520	strong similarity to long-chain-fatty-acid-CoA ligase Faa2 - Saccharomyces cerevisiae	366.53	169.77	224.33	235.20	AKL	
Pc12g07380	weak similarity to allantoate permease Dal5 - Saccharomyces cerevisiae	22.67	10.50	12.17	8.25	CHI	
Pc12g08530	strong similarity to long-chain acyl-CoA dehydrogenase like protein An13g03940 - Aspergillus niger	135.73	126.77	178.30	352.00	AKI	
Pc12g09130	similarity to hypothetical protein ncu04017.1 - Neurospora crassa	291.80	245.33	212.33	224.95	SHL	
Pc12g09740	strong similarity to fructosyl amine oxygen oxidoreductase - Aspergillus fumigatus	79.50	53.60	42.57	63.08	AKL	
Pc12g09950	alcohol oxidase aox - Penicillium chrysogenum	6.23	4.13	9.13	2.73	SRL	
Pc12g13480	similarity to protein #3345 from patent US6562958-B1 - Acinetobacter baumannii	4.40	2.93	52.57	104.65	SKL	
Pc12g14790	strong similarity to AK-toxin production protein Akt3-1 - Alternaria alternata	2.43	0.27	2.87	1.18	AKL	
Pc12g15900	hypothetical protein	1.57	1.63	2.87	2.23	ARI	
Pc13g01380	strong similarity to fructosyl amine oxygen oxidoreductase - Aspergillus fumigatus	7.57	7.00	9.13	11.18	SKL	
Pc13g01530	strong similarity to hypothetical protein Afu2g02950 - Aspergillus fumigatus	34.17	42.67	67.90	123.28	CKI	
Pc13g01660	similarity to herbicidally active polypeptide SEQ ID NO 2512 from patent WO200210210-A2 - Arabidopsis thaliana	10.33	16.87	70.13	374.68	SKL	
Pc13g01890	strong similarity to acyl-CoA synthase like protein An18g03800 - Aspergillus niger	144.97	223.63	132.00	391.13		RLSGLLGHI
Pc13g02660	strong similarity to hypothetical protein AN7738_2 - Aspergillus nidulans	6.83	3.10	6.47	9.45	AKL	
Pc13g02710	strong similarity to multifunctional beta-oxidation protein - Neurospora crassa	441.87	416.23	460.40	853.38	SKL	
Pc13g02990	strong similarity to pyruvate dehydrogenase phosphatase isoenzyme 1 PDP1 - Rattus norvegicus	231.50	238.93	200.73	179.30	SKL	
Pc13g03100	weak similarity to hypothetical protein An12g03380 - Aspergillus niger [putative sequencing error]	11.70	11.10	15.73	18.35	ARL	
Pc13g03340	strong similarity to succinate-semialdehyde dehydrogenase NAD(P)+ gabD - Escherichia coli	1.37	0.20	2.90	1.53	SKL	
Pc13g03350	strong similarity to copper amine oxidase AO-I - Aspergillus niger	15.10	20.60	22.10	18.93	CRL	
Pc13g03610	strong similarity to hypothetical protein AAF47250.1 - Drosophila melanogaster	85.20	199.30	180.13	524.75	SKI	
Pc13g03720	hypothetical protein	1.80	3.17	4.23	1.23	ARM	
Pc13g04270	strong similarity to dimethylglycine dehydrogenase precursor DMGDH - Homo sapiens	17.03	15.13	36.57	30.08	SRL	
Pc13g04370	similarity to peroxisomal membrane protein PEX17 - Yarrowia lipolytica	262.23	271.40	245.67	273.85	SKL	
Pc13g04510	strong similarity to NADP-dependent malate dehydrogenase mdh - Homo sapiens	97.87	98.20	81.33	33.35	AKL	
Pc13g05130	strong similarity to acetyl coenzyme A synthase - Tetrahymena pyriformis	64.07	60.80	39.87	56.75	SKL	
Pc13g05220	strong similarity to microsomal cytochrome like protein An05g00300 - Aspergillus niger	729.80	864.43	786.00	1126.73	AKL	
Pc13g05940	strong similarity to trifunctional protein of the beta-oxidation fox-2 - Neurospora crassa	491.57	263.90	427.53	322.68	AKL	
Pc13g06220	strong similarity to hypothetical protein brt1 - Schizophyllum commune	276.40	309.57	404.73	383.55	SRL	
Pc13g06300	strong similarity to acetamidase amdS - Aspergillus nidulans	6.63	146.67	32.77	225.40	SKI	
Pc13g06370	strong similarity to hypothetical protein 1148_scaffold_2.tfa_890cg - Fusarium graminearum	4.40	4.30	16.43	15.85	AKL	
Pc13g07280	strong similarity to hypothetical protein 1158_scaffold_2.tfa_100wg - Fusarium graminearum	19.97	6.33	22.23	4.33		KVLHLLQHA
Pc13g07430	strong similarity to hypothetical protein Afu4g07940 - Aspergillus fumigatus	57.53	37.17	106.77	226.33	SKL	
Pc13g09070	strong similarity to hypothetical protein B23L21.110 - Neurospora crassa	64.30	48.83	48.10	54.45	ARL	
Pc13g09740	strong similarity to fructosyl amine oxygen oxidoreductase - Aspergillus fumigatus	48.27	43.43	117.77	193.65	AKL	
Pc13g11580	strong similarity to hypothetical protein Afu1g12880 - Aspergillus fumigatus	120.37	326.87	112.03	458.30	ARL	
Pc13g11930	strong similarity to acyl CoA dehydrogenase aidB - Escherichia coli	1.37	2.83	51.37	108.40	SKL	
Pc13g12080	similarity to acetyl-hydrolase chnC - Acinetobacter sp.	111.07	147.07	180.30	440.23	ARL	
Pc13g12270	strong similarity to 4-coumarate-CoA ligase 4CL - Arabidopsis thaliana	92.17	100.77	57.33	133.35	AKL	
Pc13g12580	strong similarity to 8-amino-7-oxononanoate synthase/KAPA synthase BioF - Kurthia sp.	0.43	2.50	2.37	0.90	ARL	KILSQKLQF

Pc13g12620	strong similarity to spindle assembly checkpoint protein like protein An08g06660 - Aspergillus niger	105.27	90.77	66.83	63.78	ARM	
Pc13g12760	strong similarity to pyruvate decarboxylase dcpY - Aspergillus parasiticus	0.57	1.23	5.53	3.98		RLHQVGLHA
Pc13g12930	strong similarity to peroxisomal acetyl-CoA C-acyltransferase POT1 - Yarrowia lipolytica	444.73	317.80	446.53	442.33		RLSSLLSHL
Pc13g14410	strong similarity to hypothetical protein Afu7g06090 - Aspergillus fumigatus	5.47	4.70	7.07	20.60	SKL	
Pc13g14420	strong similarity to malonyl CoA synthase matB - Rhizobium trifolii	4.67	8.50	7.73	86.88	ARL	
Pc13g14580	hypothetical protein	2.30	2.00	2.43	2.58	AKL	
Pc13g15940	strong similarity to copper amine oxidase AO-I - Aspergillus niger	200.00	365.70	98.40	195.53	SHI	
Pc14g00140	strong similarity to hypothetical protein AN3880_2 - Aspergillus nidulans	69.67	126.00	163.67	95.90	ARI	
Pc14g00560	strong similarity to hypothetical protein EAA30321.1 - Neurospora crassa	36.20	39.47	34.57	53.25	AKL	
Pc14g00970	similarity to D-amino-acid oxidase DAO - Fusarium solani	2.23	0.63	2.17	3.03	SKL	
Pc14g01040	strong similarity to aldehyde dehydrogenase aldA - Aspergillus niger	1.33	1.03	0.90	1.33	ARL	
Pc14g01070	strong similarity to actVA-ORF4-like protein from patent WO9911793-A1 - Homo sapiens	15.50	8.43	12.37	10.70	SKI	
Pc14g01090	strong similarity to NADH oxidase nadA - Aspergillus parasiticus	2.13	0.23	2.13	0.78	SKL	
Pc14g01270	strong similarity to 2-hydroxyhepta-2,4-diene-1,7-dioate isomerase BH2000 - Bacillus halodurans	269.07	364.33	481.37	806.98	SKL	
Pc15g00240	weak similarity to hypothetical protein Afu6g10160 - Aspergillus fumigatus	1.27	0.43	2.40	0.70	SRI	
Pc15g00410	strong similarity to peroxisomal acetyl-CoA C-acyltransferase POT1 - Yarrowia lipolytica	185.87	149.63	163.77	271.45		RLNSVANQL
Pc15g00420	strong similarity to very long-chain fatty acyl-CoA synthase Fat1 - Saccharomyces cerevisiae	3.07	1.47	3.60	8.35	AKI	
Pc15g00830	strong similarity to hypothetical protein AN0303_2 - Aspergillus nidulans	87.37	91.40	74.53	72.90		RQVDHAAHA
Pc16g00190	strong similarity to NADH oxidase nadA - Aspergillus parasiticus	2.43	3.13	3.77	9.68	SHL	
Pc16g03770	strong similarity to conserved hypothetical protein - Brucella suis	116.23	75.23	103.23	149.05	SRL	
Pc16g03970	strong similarity to hypothetical conserved protein SPCP1E11.11 - Schizosaccharomyces pombe	130.23	124.37	119.77	49.53	SKI	
Pc16g04350	strong similarity to hypothetical protein An02g13300 - Aspergillus niger	3.53	2.33	5.70	12.73	SKL	
Pc16g05060	strong similarity to 3-hydroxy-3-methylglutaryl coenzyme A reductase HmgA - Sulfolobus solfataricus	0.23	0.33	1.90	1.68	SKI	
Pc16g05620	strong similarity to protein kinase CK2 (casein kinase II) beta subunit ckb1p - Schizosaccharomyces pombe [truncated ORF]	331.80	348.57	219.47	192.68	AKM	
Pc16g05900	similarity to salicylate hydroxylase nahG - Pseudomonas putida	0.27	0.23	0.53	0.25	ARM	
Pc16g06490	strong similarity to hypothetical protein An04g06530 - Aspergillus niger	2.33	0.63	3.87	3.43	SRL	
Pc16g07060	strong similarity to catalase C catC - Aspergillus nidulans	4.33	2.07	3.20	2.88	SRL	
Pc16g07070	strong similarity to aryl-alcohol oxidase precursor aao - Pleurotus eryngii	3.53	2.33	3.87	2.75	SKL	
Pc16g07440	strong similarity to cytoplasmic proline--tRNA ligase like protein An08g02860 - Aspergillus niger	44.50	41.83	44.73	51.28	AKL	
Pc16g10060	strong similarity to methylmalonate-semialdehyde dehydrogenase precursor MMSDH - Bos taurus [putative sequencing error]	6.97	4.93	6.13	2.75	SHL	
Pc16g10070	strong similarity to 2-nitropropane dioxygenase ncd-2 - Neurospora crassa	10.67	13.97	12.17	13.00	SKL	
Pc16g10670	strong similarity to cyclohexanone monooxygenase chnB - Acinetobacter sp.	10.80	11.73	14.17	14.35	SKL	
Pc16g11360	similarity to tripeptidylaminopeptidase Tap - Streptomyces lividans	143.60	109.83	125.30	98.03		RQVRQAHQA
Pc16g11450	strong similarity to hypothetical protease Mch5 - Saccharomyces cerevisiae	2.97	2.63	18.70	23.63	AKI	
Pc16g11790	strong similarity to fructosyl amine oxygen oxidoreductase - Aspergillus fumigatus	5.57	10.43	10.60	14.30	SKI	
Pc16g11910	similarity to hypothetical protein BAB09014.1 - Arabidopsis thaliana	5.17	2.37	5.90	6.00	SKL	
Pc16g14240	strong similarity to acetamidase amdS - Aspergillus oryzae	220.13	169.30	228.40	272.30	AKL	
Pc16g14920	similarity to zinc-finger protein like protein An12g08230 - Aspergillus niger	0.50	0.93	2.23	1.25	AKL	
Pc17g00250	hypothetical protein	25.30	18.23	21.57	17.03	SRM	
Pc17g00720	similarity to hypothetical protein AN0335_2 - Aspergillus nidulans	0.63	2.23	2.80	1.58	ARL	
Pc17g00980	hypothetical protein	na	na	na	na	CKI	
Pc18g01590	strong similarity to glycolate oxidase GOX - Mus musculus	13.97	16.50	19.80	35.85	ARL	
Pc18g01860	strong similarity to D-amino-acid oxidase DAO - Fusarium solani	10.13	8.27	6.93	6.78	SHL	
Pc18g03000	strong similarity to hypothetical protein YMR099c - Saccharomyces cerevisiae	455.80	460.13	554.63	501.10	SRL	
Pc18g03470	strong similarity to lysine aminopeptidase apsA - Aspergillus niger	436.43	450.17	305.93	236.55	SKL	
Pc18g04530	similarity to hypothetical coiled-coil protein SPAC13A11.03 - Schizosaccharomyces pombe	48.00	45.50	46.90	47.25		KQDLILTHL
Pc19g00210	strong similarity to hypothetical protein 1190_scaffold_2.tfa_540wg - Fusarium graminearum [putative pseudogene]	2.63	2.50	3.17	0.38	ARI	
Pc20g01800	strong similarity to hypothetical protein Afu7g06090 - Aspergillus fumigatus	438.57	118.13	125.37	230.88	SKL	
Pc20g01980	strong similarity to phenol 2-monooxygenase - Trichosporon beigelii	9.70	60.60	12.40	79.53	SRL	
Pc20g03400	hypothetical protein	3.17	2.33	3.77	3.25	ARL	
Pc20g03610	strong similarity to precursor of mitochondrial isocitrate dehydrogenase icdA - Aspergillus niger	867.90	1270.00	943.53	1200.78	ARL	
Pc20g03630	strong similarity to acetamidase amdS - Aspergillus nidulans	4.33	2.90	2.37	2.83	ARL	
Pc20g04300	strong similarity to hypothetical protein An16g02770 - Aspergillus niger	50.20	83.93	237.90	507.25	ARL	
Pc20g06660	strong similarity to D-amino-acid oxidase DAO - Fusarium solani	29.83	24.80	63.20	79.48	SRL	

Pc20g08300	strong similarity to hypothetical protein 12F11.200 - <i>Neurospora crassa</i>	51.20	76.80	72.30	150.13	SKI	
Pc20g09430	similarity to hypothetical ureidoglycolate hydrolase SPAC19G12.04 - <i>Schizosaccharomyces pombe</i>	19.73	31.90	40.17	51.18	AKL	
Pc20g10430	strong similarity to D-mandelate dehydrogenase - <i>Rhodotorula graminis</i>	1.77	2.87	4.77	2.58	SKL	
Pc20g10520	similarity to hypothetical protein An12g07270 - <i>Aspergillus niger</i>	11.90	22.20	13.67	29.63	AKM	
Pc20g10700	strong similarity to hypothetical protein 1143_scaffold_2.tfa_210cg - <i>Fusarium graminearum</i>	1.43	3.27	3.23	4.43	SKI	
Pc20g11660	strong similarity to mitochondrial mRNA processing protein Pet127 - <i>Saccharomyces cerevisiae</i>	71.93	61.13	51.40	26.65	CKM	
Pc20g13550	strong similarity to malate synthase acuE - <i>Aspergillus nidulans</i>	164.57	123.83	397.17	188.58	AKL	
Pc20g14920	similarity to hypothetical oxidoreductase PA5309 - <i>Pseudomonas aeruginosa</i>	11.73	39.70	29.90	96.10	ARL	
Pc20g15650	strong similarity to hypothetical protein AN5853_2 - <i>Aspergillus nidulans</i>	6.77	2.00	11.70	41.18	ARL	
Pc20g15660	strong similarity to sterol carrier protein-X/sterol carrier protein-2 SCP2 - <i>Homo sapiens</i>	6.03	6.27	23.63	107.70	ARI	
Pc21g00200	hypothetical protein	na	na	na	na		KLRCISCHL
Pc21g00230	hypothetical protein	na	na	na	na	CKI	
Pc21g00970	strong similarity to hex1 - <i>Aspergillus nidulans</i>	2685.47	2952.60	2926.47	2448.45	SRL	
Pc21g01040	similarity to hypothetical protein AN7405_2 - <i>Aspergillus nidulans</i>	89.63	86.97	83.97	77.23	SRM	
Pc21g01650	strong similarity to hypothetical conserved protein PA5185 - <i>Pseudomonas aeruginosa</i>	97.43	132.47	117.50	237.35	AKL	
Pc21g02970	strong similarity to phenol 2-monooxygenase - <i>Trichosporon beigelii</i>	64.27	80.87	73.63	104.65	SRL	
Pc21g03400	strong similarity to triose-phosphate-isomerase tpiA from patent WO8704464-A - <i>Aspergillus niger</i>	1488.47	1672.60	2271.77	1813.75	ARI	
Pc21g03940	strong similarity to transposase Tan1 - <i>Aspergillus niger</i> [putative pseudogene]	2.90	1.47	4.43	5.00	ARI	
Pc21g04900	strong similarity to hypothetical protein AN4261_2 - <i>Aspergillus nidulans</i>	89.97	83.13	83.87	126.78	SKL	
Pc21g05470	strong similarity to mannase man1 - <i>Aspergillus aculeatus</i>	6.40	4.53	6.27	9.05	SKI	
Pc21g05490	strong similarity to sn-glycerol-3-phosphate acyltransferase like protein An18g01960 - <i>Aspergillus niger</i>	189.90	157.30	188.90	201.75	AKL	
Pc21g05590	strong similarity to D-amino-acid oxidase DAO - <i>Fusarium solani</i>	304.87	383.33	278.93	372.93	AKL	
Pc21g05690	weak similarity to hypothetical protein An09g06720 - <i>Aspergillus niger</i>	1.90	0.93	3.13	2.18	SHM	
Pc21g06590	strong similarity to hypothetical protein Afu8g02420 - <i>Aspergillus fumigatus</i>	2.87	1.90	6.93	9.45	SRL	
Pc21g06860	strong similarity to endopeptidase La 2 lonD - <i>Myxococcus xanthus</i>	316.17	291.53	356.00	378.08	SRL	
Pc21g07100	similarity to transposase Taf1 - <i>Aspergillus fumigatus</i> [putative pseudogene]	2.70	1.20	0.73	0.73	SKI	
Pc21g07210	weak similarity to S-layer protein CipA - <i>Clostridium thermocellum</i>	348.73	315.87	215.77	190.35	SKL	
Pc21g07490	strong similarity to hypothetical protein AN0070_2 - <i>Aspergillus nidulans</i>	65.43	62.93	62.67	71.93	SRI	
Pc21g08790	strong similarity to NADP-dependent leukotriene B4 12-hydroxydehydrogenase like protein An13g01120 - <i>Aspergillus niger</i>	58.93	45.00	86.93	99.93	AKL	
Pc21g09440	strong similarity to hypothetical protein AN5270_2 - <i>Aspergillus nidulans</i>	1.10	2.87	5.53	4.23	SKL	
Pc21g09460	strong similarity to propionyl-CoA carboxylase, beta subunit - <i>Mycobacterium tuberculosis</i>	1.40	0.63	1.63	0.20	ARI	
Pc21g09470	strong similarity to feruloyl-CoA synthase like protein An06g01320 - <i>Aspergillus niger</i>	0.23	0.20	0.80	0.33	AKL	
Pc21g09480	strong similarity to 3-hydroxy-3-methylglutaryl CoA lyase HMG-CoA lyase - <i>Rattus norvegicus</i>	4.87	1.63	5.23	5.33	SKL	
Pc21g09780	strong similarity to fructosyl amine oxygen oxidoreductase - <i>Aspergillus fumigatus</i>	8.17	4.53	30.70	43.25	SRL	
Pc21g10640	strong similarity to cadmium resistance protein - <i>Saccharomyces cerevisiae</i>	374.47	342.17	413.80	371.18	SKI	
Pc21g11080	strong similarity to hypothetical protein AN1437_2 - <i>Aspergillus nidulans</i>	93.57	64.93	52.63	32.83	SRM	
Pc21g11670	strong similarity to quinone reductase like protein An12g00170 - <i>Aspergillus niger</i>	84.90	77.50	102.33	139.55	SKL	
Pc21g11780	strong similarity to 3-dehydroshikimate dehydratase qa-4 - <i>Neurospora crassa</i>	3.07	4.57	11.43	32.60	SRI	
Pc21g12260	strong similarity to alcohol dehydrogenase C ADHC - <i>Mycobacterium smegmatis</i>	20.73	12.07	22.40	23.20	AKL	
Pc21g13110	strong similarity to polyprotein of retrotransposon REAL pol - <i>Alternaria alternata</i> [putative pseudogene]	0.17	0.40	0.63	1.40	SRI	
Pc21g13270	strong similarity to hypothetical protein AN3770_2 - <i>Aspergillus nidulans</i>	37.13	37.77	40.27	52.38	SRM	
Pc21g14590	strong similarity to protease involved in a-factor processing Ste23 - <i>Saccharomyces cerevisiae</i>	299.93	303.43	362.53	285.30	AKL	
Pc21g14640	strong similarity to hypothetical protein YDR196c - <i>Saccharomyces cerevisiae</i>	53.07	62.70	82.40	175.88	AKL	
Pc21g15970	strong similarity to cytosolic exopolyphosphatase Ppx1 - <i>Saccharomyces cerevisiae</i>	31.23	29.47	54.83	83.50	AKL	
Pc21g16050	strong similarity to hypothetical protein YGL067w - <i>Saccharomyces cerevisiae</i>	76.30	122.50	93.77	154.23	SKI	
Pc21g16080	strong similarity to hypothetical protein encoded by gene B18D24.50 - <i>Neurospora crassa</i>	184.10	242.00	576.63	431.73	AHI	
Pc21g16580	hypothetical protein	6.00	3.23	5.40	4.50	AKI	
Pc21g16710	strong similarity to hypothetical protein AACM01000094_6 - <i>Fusarium graminearum</i>	22.20	72.73	14.67	62.95	ARL	
Pc21g17590	strong similarity to acyl-CoA dehydrogenase like protein An17g01150 - <i>Aspergillus niger</i>	529.80	553.13	838.73	1248.65	SHL	
Pc21g20650	strong similarity to 4-coumarate-CoA ligase 4CL - <i>Arabidopsis thaliana</i>	72.60	62.87	52.57	123.08	ARL	RLQALSKQ
Pc21g21120	strong similarity to monoamine oxidase N mao-N - <i>Aspergillus niger</i>	0.83	11.57	0.53	14.25	SHL	
Pc21g21140	similarity to hypothetical protein - <i>Gloeobacter violaceus</i>	2.63	1.00	3.37	5.15	AHL	
Pc21g21370	acyl-coenzyme A:isopenicillin N acyltransferase (acyltransferase) AAT/PenDE - <i>Penicillium chrysogenum</i>	1031.33	1827.00	1993.70	2714.55	ARL	
Pc21g21410	strong similarity to hypothetical protein ncu05765.1 - <i>Neurospora crassa</i>	2.63	1.70	5.40	6.30	SRM	
Pc21g21810	strong similarity to levodione reductase lvr - <i>Corynebacterium aquaticum</i>	79.30	82.27	154.37	241.25		
Pc21g22010	strong similarity to 4-coumarate-CoA ligase 4CL - <i>Arabidopsis thaliana</i>	50.77	45.30	22.03	11.73	SKL	
Pc21g23700	strong similarity to long-chain-fatty-acid alcohol oxidase fao1 - <i>Candida cloacae</i>	54.73	66.50	40.03	55.33	ARL	
Pc22g00060	similarity to fructosyl amine oxygen oxidoreductase - <i>Aspergillus fumigatus</i>	93.93	177.60	137.93	334.18	AKL	
Pc22g00300	strong similarity to flavocytochrome b2 L-lactate dehydrogenase CYB2 - <i>Pichia anomala</i>	8.87	7.37	9.57	4.35	SKL	

Pc22g00460	strong similarity to hypothetical protein An06g01070 - Aspergillus niger	62.90	67.23	227.67	87.93	ARI	
Pc22g00860	strong similarity to cytochrome-c peroxidase precursor Ccp1 - Saccharomyces cerevisiae	62.63	3.03	5.67	1.98	ARL	
Pc22g01300	similarity to lipid transfer protein POX18 - Candida tropicalis	407.10	286.73	628.57	403.55	AKL	
Pc22g01600	strong similarity to histidine triad protein Hnt1 - Saccharomyces cerevisiae	882.90	1065.03	1135.30	1128.53	SKM	
Pc22g02570	hypothetical protein	6.17	6.83	5.93	7.35	AHI	
Pc22g02620	strong similarity to monoglyceride lipase mgII - Mus musculus	71.67	119.07	89.37	238.30	AKL	
Pc22g03680	strong similarity to NADPH-dependent beta-ketoacyl reductase RhlG - Pseudomonas aeruginosa	161.83	130.50	370.53	460.73	AKL	
Pc22g05640	similarity to hypothetical protein required for biosynthesis of the host-specific AK-toxin Akt2 - Alternaria alternata	13.07	18.37	13.57	29.08	SKL	
Pc22g06120	strong similarity to SR-protein-specific kinase SRPK2 - Mus musculus	460.47	397.50	324.80	220.25	SHM	
Pc22g06820	strong similarity to peroxisomal acetyl-CoA C-acyltransferase POT1 - Yarrowia lipolytica	475.30	194.27	244.33	157.40		RLSQVSSHf
Pc22g07740	strong similarity to acyl-CoA oxydase tylP - Streptomyces fradiae	2.73	0.87	2.30	2.33	AKL	
Pc22g08120	similarity to hypothetical meta-cleavage compound hydrolase gene ren71 - Streptomyces aureofaciens [putative sequencing error]	1.30	1.13	8.80	11.53	ARI	
Pc22g10030	strong similarity to betaine-aldehyde dehydrogenase betB - Escherichia coli	373.37	568.73	212.27	232.75	AKL	
Pc22g11470	strong similarity to L-lactate dehydrogenase precursor Cyb2 - Saccharomyces cerevisiae	142.77	177.83	147.63	271.18	ARL	
Pc22g13390	strong similarity to alcohol dehydrogenase like protein An02g02060 - Aspergillus niger	258.07	301.53	529.07	917.10	SKL	
Pc22g13680	strong similarity to hypothetical protein Afu5g03740 - Aspergillus fumigatus	17.27	13.67	12.30	15.40	SKI	
Pc22g14270	strong similarity to hypothetical protein Afu5g09600 - Aspergillus fumigatus	231.20	118.90	318.57	333.53	SKL	
Pc22g14900	phenylacetyl-CoA ligase pclA - Penicillium chrysogenum	108.53	139.97	222.50	381.63	SKI	
Pc22g15030	strong similarity to peroxisomal 2,4-dienoyl-CoA reductase involved in sporulation Sps19 - Saccharomyces cerevisiae	428.20	555.93	388.37	813.73	SKL	
Pc22g15400	similarity to hypothetical protein required for biosynthesis of the host-specific AK-toxin Akt2 - Alternaria alternata	8.67	0.97	0.73	0.58	SKI	
Pc22g17230	strong similarity to aldehyde dehydrogenase aldA - Aspergillus niger	9.00	6.43	14.70	11.60	SKL	
Pc22g18190	strong similarity to L-lactate dehydrogenase precursor Cyb2 - Saccharomyces cerevisiae	39.83	59.00	70.93	113.65	SKL	
Pc22g18380	strong similarity to salicylate hydroxylase nahG - Pseudomonas putida	14.47	40.33	38.87	69.70	SRL	
Pc22g18490	strong similarity to hypothetical protein T22K18.2 - Arabidopsis thaliana	107.20	136.47	111.80	189.13	CRM	
Pc22g18850	similarity to secretory polypeptide SPTM SEQ ID NO 850 from patent WO20028376-A2 - Homo sapiens	113.83	109.43	122.23	117.63	ARL	
Pc22g19270	strong similarity to glycolate oxidase GOX - Mus musculus	0.13	1.60	0.60	0.23	SKL	
Pc22g19440	strong similarity to cytosolic aspartate transaminase Aat2 - Saccharomyces cerevisiae	626.00	780.57	930.77	1212.05	AKL	
Pc22g19490	strong similarity to trans-2-enoyl-ACP reductase like protein An16g05340 - Aspergillus niger	923.17	504.00	1072.27	956.15	SKL	
Pc22g20270	strong similarity to 4-coumarate-CoA ligase 4CL - Arabidopsis thaliana	111.83	173.90	186.27	458.80	AKL	
Pc22g20320	strong similarity to myo-inositol 2-dehydrogenase yisS - Bacillus subtilis	75.60	63.33	53.03	66.13	ARL	
Pc22g20370	strong similarity to carnitine racemase like protein An03g03550 - Aspergillus niger	0.67	1.60	4.93	43.33	SKL	
Pc22g20380	strong similarity to aerobactin biosynthesis protein iucB - Escherichia coli	6.47	15.03	32.73	90.15	AKL	
Pc22g20960	strong similarity to urate oxidase uaz - Aspergillus flavus	107.13	97.33	318.03	96.15	AKL	
Pc22g21240	strong similarity to catalase cta1p - Schizosaccharomyces pombe	54.93	61.93	34.00	35.80	ARL	
Pc22g21670	strong similarity to endocytosis protein Ede1 - Saccharomyces cerevisiae	417.50	370.63	304.93	205.28		RVFFQLFQA
Pc22g22110	strong similarity to glycerol-3-phosphate dehydrogenase (NAD+) precursor Gpd1 - Saccharomyces cerevisiae	316.37	311.97	248.20	201.75	ARL	
Pc22g22390	strong similarity to AK-toxin production protein Akt3-1 - Alternaria alternata	28.37	30.93	51.27	141.38	SKL	
Pc22g23350	strong similarity to fructosyl amine oxygen oxidoreductase - Aspergillus fumigatus	0.50	1.80	1.50	3.53	AKL	
Pc22g23700	weak similarity to fructosyl amine oxygen oxidoreductase - Aspergillus fumigatus	12.37	15.13	32.37	52.00	SRL	
Pc22g24630	similarity to protein involved in cephalosporin C biosynthesis like protein An09g06420 - Aspergillus niger	3.77	5.43	6.03	7.00	AKI	
Pc22g24780	strong similarity to 4-coumarate-CoA ligase 4CL - Arabidopsis thaliana	11.60	44.10	18.17	252.33	AKL	
Pc22g25120	strong similarity to hypothetical protein 1141_scaffold_2.tfa_360cg - Fusarium graminearum	12.03	20.07	14.80	38.23	SKL	
Pc22g25150	strong similarity to acyl CoA dehydrogenase aidB - Escherichia coli	46.17	17.53	96.90	189.28	SKL	
Pc22g26560	hypothetical protein	na	na	na	na		KLRCISCHL
Pc22g26600	hypothetical protein	na	na	na	na	CKI	
Pc23g00390	similarity to hypothetical protein ncu04490.1 - Neurospora crassa	36.70	42.07	51.67	34.03	AKL	
Pc24g00180	hypothetical protein [truncated ORF]	na	na	na	na	CKI	
Pc24g01320	similarity to hypothetical S-layer protein - Clostridium thermocellum	231.23	353.20	314.27	357.38	SRM	
Pc24g02680	strong similarity to copia-like retrotransposable element - Arabidopsis thaliana	4.30	2.20	2.30	3.90	CRL	
Pc32g00010	strong similarity to protein encoded by ORF1 of transposon Ant1 - Aspergillus niger	23.17	30.97	13.63	2.55	CHL	

Supplementary Table 8: Summary of the global transcriptional response to phenylacetic acid in chemostat cultures of *P. chrysogenum*. Transcript levels of strains_Wisconsin54-1255 and DS17690 were analysed in triplicate glucose-limited chemostat cultures in the absence and presence of phenylacetic acid (PAA); **A.** Numbers of genes whose transcript levels were below the detection limit in each situation. Transcript levels of 3970 genes were below detection limit in all four situation; **B.** Pairwise comparisons, showing the numbers of genes that showed a significantly different transcript level between the two strains or in response to PAA addition. Transcript levels of 2470 genes were significantly different in at least one of the four situations.

A.

	Below detection limit
DS17690 - PAA	4552
DS17690+ PAA	5054
Wis 54-1255 - PAA	4755
Wis 54-1255 + PAA	5213

B.

	Higher transcript level	Lower transcript level
DS17690 + PAA vs. DS1760 - PAA	443	411
Wis54-1255 + PAA vs. Wis54-1255 - PAA	252	338
DS17690 + PAA vs. Wis54-1255 + PAA	861	744
DS17690 - PAA vs. Wis54-1255 - PAA	492	549

Supplementary Table 9. K-mean cluster 1

@=Values given are the average of three independent experiments

orf code	Description of putative <i>P. chrysogenum</i> ORF	Best blast homolog to putative <i>P. chrysogenum</i> ORF			Average transcript levels @				FunCat (auto)	
		p-value	Gene code	description	WIS - PAA	WIS + PAA	DS - PAA	DS + PAA	pval	category
Pc06g01600	strong similarity to FAD dependent L-sorbose dehydrogenase SDH - Gluconce	2e-69	GO0622_2	product: "L-sorbose dehydrogenase	41.2	2373.4	20.6	2223.1	0.0	02.11 electron transport and membrane-associated ener
Pc12g02570	strong similarity to cytochrome P450 monooxygenase TR111 - Fusarium spor	4e-84	AF155773_13	gene: "FUM12"; product: "Fum12p";	12.0	295.5	12.0	922.2	1e-100	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc12g07980	similarity to benzoylformate decarboxylase - Pseudomonas putida	1e-129	NCB2G14_13	gene: "B2G14.130"; product: "relate	29.2	424.1	63.5	1227.0	1e-136	01.01.01 amino acid biosynthesis
Pc12g09020	strong similarity to maleylacetate isomerase maiA - Aspergillus nidulans	2e-76	EN1836_1	gene: "maiA"; product: "maleylaceto	47.0	3267.8	106.3	3479.8	5e-57	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc12g09030	strong similarity to fumarylacetoacetase - Homo sapiens	0.0	ENFUHY_1	gene: "fahA"; product: "fumarylaceto	138.0	3786.2	279.5	3905.8	0.0	01 METABOLISM
Pc12g09040	strong similarity to 3,4-dihydroxyphenylacetate 2,3-dioxygenase hmgA - Aspe	0.0	A57435	3,4-dihydroxyphenylacetate 2,3-diox	135.0	3551.7	202.7	3736.3	0.0	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc12g13630	strong similarity to allantoinase permease Dal5 - Saccharomyces cerevisiae	0.0	BX842634_3	gene: "B16B8.030"; product: "relate	12.0	85.7	12.0	140.9	5e-64	01.05.07 C-compound, carbohydrate transport
Pc12g13800	strong similarity to allantoinase permease Dal5 - Saccharomyces cerevisiae	5e-80	T41604	probable membrane transport protei	12.0	482.7	12.0	470.8	2e-53	01.04.07 phosphate transport
Pc13g06300	strong similarity to acetamidase amdS - Aspergillus nidulans	1e-126	T41382	acetamidase - fission yeast (Schizo	12.0	146.7	32.8	225.4	7e-95	01.01.10 amino acid degradation (catabolism)
Pc13g10030	strong similarity to high-affinity nicotinic acid permease Tna1 - Saccharomyc	1e-103	NC64C2_19	gene: "64C2.200"; product: "related	22.9	487.1	125.0	1344.0	2e-53	01.04.07 phosphate transport
Pc13g10900	strong similarity to high-affinity nicotinic acid permease Tna1 - Saccharomyc	1e-180	BX649605_57	product: "transporter, putative"; Aspe	22.9	487.1	125.0	1344.0	1e-56	01.04.07 phosphate transport
Pc16g01770	strong similarity to phenylacetate hydroxylase pahA - Penicillium chrysogenum	1e-112	AF057559_1	gene: "pahA"; product: "phenylacetat	12.0	2661.0	32.8	2533.9	1e-117	01.01.01 amino acid biosynthesis
Pc16g07590	strong similarity to acetamidase amdS - Aspergillus oryzae	0.0	AF349510_1	gene: "gmdB"; product: "general ami	66.0	2790.0	76.8	3542.9	1e-63	01.01.10 amino acid degradation (catabolism)
Pc18g01290	strong similarity to dihydroserine transporter Dtr1 - Saccharomyces cerevisiae	3e-71	AF141925_12	product: "unknown"; Aspergillus terr	20.4	175.2	42.8	224.3	6e-55	01.05.07 C-compound, carbohydrate transport
Pc20g02710	weak similarity to DOPA-dioxygenase dodA - Amanita muscaria	3e-51	BX897679_2	gene: "B2C22.020"; product: "conse	13.9	89.1	31.3	373.8	0	0
Pc20g14540	strong similarity to alcohol dehydrogenase like protein An15g07870 - Aspergil	1e-27	AP005953_4	gene: "blr4874"; Bradyrhizobium jap	98.2	868.9	127.6	798.8	9e-77	01.05 C-compound and carbohydrate metabolism
Pc21g04400	strong similarity to mandelate like protein An13g00920 - Aspergillus niger	3.00E-90	AB0793	probable MR-MLE-family protein ST	12.0	239.3	12.0	309.3	6e-75	01 METABOLISM
Pc21g12990	strong similarity to polyamine transport protein Tpo1 - Saccharomyces cerevi	1e-149	BX908812_25	gene: "G17A4.250"; product: "relate	12.0	76.6	12.0	109.1	8e-54	01.05.07 C-compound, carbohydrate transport
Pc21g14280	phenylacetate hydroxylase pahA - Penicillium chrysogenum	0.0	AF056978_1	gene: "pahA"; product: "phenylacetat	12.0	93.0	40.2	202.2	1e-111	01.01.10 amino acid degradation (catabolism)
Pc21g19470	strong similarity to mitochondrial succinate-fumarate transporter Sfc1 - Sacch	1e-125	BX897673_8	gene: "B13M15.080"; product: "prob	25.0	344.1	48.3	361.2	3e-77	01.05.04 regulation of C-compound and carbohydrate util
Pc21g21750	strong similarity to flavin-containing monooxygenase like protein An13g02690	1e-45	SPBP16F5_8	gene: "SPBP16F5.08c"; S. pombe c	25.0	344.1	48.3	361.2	1e-112	01.02.01.09.99 other catabolism of nitrogenous compou
Pc22g07320	strong similarity to mycosporosate synthase like protein An02g14970 - Aspergil	2e-91	NCB8P8_9	gene: "B8P8.090"; product: "probal	16.8	94.7	47.6	647.5	1e-80	01 METABOLISM
Pc22g07360	strong similarity to 1,2-dichlorophenol hydroxylase tfdB - Pseudomonas putid	1e-124	AY078159_14	gene: "tfdB"; product: "dichlorophen	33.2	476.9	57.7	501.2	1e-78	01.05.01.01.09 aerobic aromate catabolism
Pc22g07410	strong similarity to trihydroxytoluene oxygenase dntD - Burkholderia cepacia	2e-45	AF076848_1	gene: "dntD"; product: "trihydroxytol	12.0	97.8	12.0	199.8	0	0
Pc22g24780	strong similarity to 4-coumarate-CoA ligase 4CL - Arabidopsis thaliana	1e-98	AY254381_1	Emericella nidulans adenylate-formir	110.2	558.2	94.8	818.2	1e-128	01 METABOLISM
Pc22g24860	strong similarity to aldehyde dehydrogenase aldA - Aspergillus niger	1e-131	NCB11B23_10	gene: "B11B23.110"; product: "prob	12.0	44.1	18.2	252.3	0.0	01.05 C-compound and carbohydrate metabolism
Pc20g02930	strong similarity to D-lactate dehydrogenase ald - Kluyveromyces lactis	1e-141	BX842682_10	gene: "B13C5.100"; product: "probal	12.0	968.8	48.3	1655.9	1e-157	01.05.01 C-compound and carbohydrate utilization
Pc16g02700	weak similarity to cutinase transcription factor beta CTF1b - Fusarium solani	7e-11	T38690	probable regulatory protein - fission	44.3	308.8	71.3	526.7	4e-77	01.01.04 regulation of amino acid metabolism
Pc16g02690	strong similarity to benomyl methotrexate resistance protein MDR1 - Candida	1e-78	BX649607_11	gene: "mdR"; product: "mfs-family m	12.0	97.0	20.8	129.6	2e-48	01.04.07 phosphate transport
Pc21g01300	strong similarity to membrane protein Tpo2 - Saccharomyces cerevisiae	0.0	ACH47683_1	gene: "celT"; product: "multidrug res	12.0	280.7	12.0	910.0	9e-67	01.05.07 C-compound, carbohydrate transport
Pc21g09220	strong similarity to fluconazole resistance transporter Flr1 - Saccharomyces c	4e-90	BX649607_11	gene: "mdR"; product: "mfs-family m	12.0	81.1	12.0	591.1	4e-51	01.05.04 regulation of C-compound and carbohydrate util
Pc21g14210	similarity to polyamine transport protein Tpo3 - Saccharomyces cerevisiae	2e-23	T41018	probable membrane transporter - fisi	15.2	402.4	158.4	1118.5	7e-55	01.04.07 phosphate transport
Pc22g14600	strong similarity to ATP-binding cassette multidrug transport protein atrB - Asj	0.0	AB028872_1	gene: "BMR1"; product: "BMR1"; Bi	12.0	272.8	12.0	111.2	0.0	01.06.13 lipid and fatty-acid transport
Pc22g20580	strong similarity to multidrug resistance protein Qdr1 - Saccharomyces cerevi	9e-67	S49888	probable membrane protein YIL121v	35.0	399.5	176.0	567.2	7e-58	01.05.07 C-compound, carbohydrate transport
Pc12g13810	strong similarity to methicillin resistance gene HmrA - Staphylococcus aureus	3e-96	NCB14A21_24	gene: "B14A21.240"; product: "relate	12.2	140.9	52.5	163.4	9e-87	01.01 amino acid metabolism
Pc22g06840	similarity to arylamine N-acetyltransferase - Gallus gallus	2e-07	AY228175_2	product: "putative arylamine N-acety	31.1	2001.2	98.9	2710.2	9e-82	01 METABOLISM
Pc22g08750	strong similarity to mitomycin C translocase mct - Streptomyces lavendulae	1e-44	BX908808_43	gene: "G21B4.430"; product: "relate	19.6	150.7	33.6	345.9	2e-48	01.05.04 regulation of C-compound and carbohydrate util
Pc22g08950	strong similarity to cytochrome P450 pisatin demethylase PDAT9 - Nectria h	4e-62	S45583	pisatin demethylase (EC 1.14.-.-) cyt	12.0	78.5	12.0	120.5	1e-105	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc12g02210	similarity to hypothetical protein contig_1_67_scaffold_4.tfa_580cg - Aspergil	3e-09	H95268	hypothetical protein SMO112 [impo	30.7	554.9	52.7	547.9	0	0
Pc12g02220	strong similarity to hypothetical protein mg00678.1 - Magnaporthe grisea	1e-34	T41554	hypothetical protein SPCC70.08c - fi	12.0	1051.1	21.5	864.5	1e-35	01.07.01 biosynthesis of vitamins, cofactors, and prosthe
Pc13g06330	similarity to hypothetical membrane protein YOL119c - Saccharomyces cerev	3e-51	NCB23B10_3	gene: "B23B10.030"; product: "relate	12.0	266.8	130.3	724.7	1e-44	01.04.07 phosphate transport
Pc16g09960	strong similarity to hypothetical protein contig_1_43_scaffold_2.tfa_610cg - A	6e-48	BX842647_291	product: "Major facilitator family tran	12.0	159.7	26.4	355.4	9e-45	01.04.07 phosphate transport
Pc16g13530	strong similarity to hypothetical protein contig_1_130_scaffold_10.tfa_180cg	0	0	0	12.0	63.0	12.0	90.8	0	0
Pc19g00540	similarity to membrane steroid hormone-binding protein MSBP - Bos taurus	4e-25	S65181	hypothetical protein YPL170w - yeas	12.0	223.0	12.0	175.2	8e-47	04.05.01.04 transcriptional control
Pc22g07140	strong similarity to hypothetical protein An04g06070 - Aspergillus niger	0	0	0	12.0	162.0	12.0	217.4	0	0
Pc22g07310	strong similarity to hypothetical protein binA - Aspergillus nidulans	5e-21	ANI011295_1	gene: "binA"; product: "hypothetical	237.4	3840.0	338.7	3996.3	0	0
Pc22g07350	strong similarity to hypothetical protein SPCC757.02c - Schizosaccharomyc	4e-49	T41593	hypothetical protein SPCC757.02c -	12.0	206.7	22.8	184.6	1e-137	11.01 stress response
Pc22g07370	strong similarity to hypothetical protein contig_1_168_scaffold_14.tfa_410wg	7e-30	NCB14A6_3	gene: "B14A6.030"; product: "probal	12.0	70.6	12.0	234.1	1e-108	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc22g07400	strong similarity to hypothetical 2-hydroxyhepta-2,4-diene-1,7-dioate isomera	2.00E-51	BX897674_13	gene: "B2N18.140"; product: "conse	12.0	180.5	12.0	197.8	2e-85	01 METABOLISM
Pc22g09710	weak similarity to hypothetical protein PA1213 - Pseudomonas aeruginosa	1e-160	NCG15G9_2	gene: "G15G9.020"; product: "conse	47.5	1043.5	36.7	1180.6	0	0
Pc22g15280	hypothetical protein	0	0	0	162.8	1715.7	508.0	2631.8	0	0
Pc22g16820	weak similarity to hypothetical transcription regulator SPBC530.05 - Schizosa	8.00E-14	T40524	hypothetical protein SPBC530.08 - fi	12.0	222.6	12.0	131.7	2e-61	01.01.04 regulation of amino acid metabolism
Pc22g22960	strong similarity to hypothetical protein B2J23.80 - Neurospora crassa	6e-91	NCB2J23_8	gene: "B2J23.080"; product: "conser	155.8	898.3	177.8	1502.1	4e-61	01.01.10 amino acid degradation (catabolism)

Supplementary Table 10. K-mean cluster 2

@=Values given are the average of three independent experiments

orf code	Description of putative <i>P. chrysogenum</i> ORF	Best blast homolog to putative <i>P. chrysogenum</i> ORF		Average transcript levels @				FunCat (auto)	
		p-value	Gene code description	WIS - PAA	WIS + PAA	DS - PAA	DS + PAA	pval	category
Pc06g00650	strong similarity to hypothetical protein 1147_scaf7e-43	0	AP050525_101 product: 'putative hydrolase'; Streptomyces avermitilis genomic DNA,	222.1	488.9	431.0	1108.5	0	
Pc06g01070	strong similarity to membrane protein Tpo2 - Sacc 1e-122	0	NCB19C19_12 gene: 'B19C19.140'; product: 'related to fluconazole resistance protein	12.0	57.7	25.5	62.0	5e-62	01.05.07 C-compound, carbohydrate transport
Pc06g01350	strong similarity to bifunctional xylosidase-arabinos 3e-26	0	AE016937_194 product: 'periplasmic beta-glucosidase precursor'; Bacteroides thetaiota	14.1	48.3	14.9	22.1	1e-117	01.05.01.01 C-compound, carbohydrate catabolism
Pc06g01360	similarity to bifunctional xylosidase-arabinosidase 1e-17	0	BFAF6658_1 product: 'beta-glucosidase'; Bacteroides fragilis beta-glucosidase gene	15.1	64.7	12.9	25.8	7e-41	01.05.01.01 C-compound, carbohydrate catabolism
Pc06g01370	strong similarity to hypothetical membrane protein 1e-139	0	AY236409_1 gene: 'mct'; product: 'monocarboxylate transporter-like protein'; Emer	17.4	57.7	12.0	23.0	1e-42	01.04.07 phosphate transport
Pc08g00080	weak similarity to scytalone dehydratase 1 - Pyricularia	0	AF029913_4 gene: 'BGL1'; product: 'beta glucosidase homolog'; Cochliobolus heter	62.9	517.1	384.5	530.1	0	
Pc12g03690	strong similarity to hypothetical protein contig_1_11e-94	0	A78767_1 unnamed ORF; Sequence 1 from Patent EP0565172.	70.7	455.4	151.6	108.6	6e-64	02.11 electron transport and membrane-associated energy conservation
Pc12g06820	strong similarity to hypothetical protein BAA10688 8e-25	0	S76996 hypothetical protein - Synechocystis sp. (strain PCC 6803)	49.5	73.2	24.1	56.5	3e-38	04.05.01.04 transcriptional control
Pc12g09010	weak similarity to transcription factor Aro80 - Sacc 3e-27	0	S69704 hypothetical protein YDR421w - yeast (Saccharomyces cerevisiae)	24.0	83.2	49.3	128.8	1e-118	04.05.01.04 transcriptional control
Pc12g10830	strong similarity to cytochrome b5 - Morielia alp 4e-38	0	NCB23L21_12 gene: 'B23L21.190'; product: 'probable cytochrome b5'; Neurospora crassa	819.9	2287.4	816.9	2887.2	1e-30	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc12g10870	strong similarity to lanosterol 14 alpha-demethylase 0.0	0	AB030178_1 gene: 'PDCYP51'; product: 'cytochrome P-450 14DM'; Penicillium dig	136.6	252.5	86.2	188.2	1e-106	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc12g11110	strong similarity to beta-glucosidase 1 bgl1 - Aspe 0.0	0	AF029913_4 gene: 'BGL1'; product: 'beta glucosidase homolog'; Cochliobolus heter	130.1	96.8	60.4	140.2	0.0	01.05.01.01 C-compound, carbohydrate catabolism
Pc12g11140	strong similarity to hypothetical protein SC8D11.1 3e-32	0	SC0939132_1 gene: 'SC07720'; 'SC8D11.11'; product: 'hypothetical protein SC8D1	12.0	69.0	13.3	87.7	0	
Pc12g11950	strong similarity to hypothetical protein SPAC1102.2e-88	0	AE016865_237 product: 'conserved hypothetical protein'; Pseudomonas syringae pv. l	28.8	80.8	67.5	127.7	2e-76	99 UNCLASSIFIED PROTEINS
Pc12g11990	strong similarity to fluconazole resistance transpor 3e-79	0	UMLPLOC_5 gene: 'pig6'; product: 'related to multidrug resistant protein'; Ustilago	54.3	172.2	63.1	214.5	1e-52	01.04.07 phosphate transport
Pc12g12000	similarity to zinc-finger transcription factor amdA - 6e-11	0	S61908 DNA-binding protein amdA - Emericella nidulans	35.5	74.4	40.1	53.7	3e-87	04 TRANSCRIPTION
Pc12g12010	weak similarity to hypothetical mlf5689 - Mesorhiz 2e-23	0	AP006569_36 Gloeobacter violaceus PCC 7421 DNA, complete genome, section 2/11	274.3	751.7	737.6	1366.3	0	
Pc12g12960	weak similarity to transcription factor Arg81 - Sacc	0	0	31.2	59.0	46.5	114.3	0	
Pc12g13110	strong similarity to hypothetical protein SPAC227.2e-35	0	NCB7H23_12 gene: 'B7H23.120'; product: 'conserved hypothetical protein'; Neuros	26.8	53.0	21.7	54.0	7e-44	01 METABOLISM
Pc12g13590	similarity to acid phosphatase PHO2 - Yarrowia lip 2e-22	0	S19993 acid phosphatase (EC 3.1.3.2) - yeast (Yarrowia lipolytica)	12.0	20.8	12.0	43.2	0	
Pc12g13640	strong similarity to transcription conserved protein 1e-142	0	BX842634_4 gene: 'B1688.040'; product: 'conserved hypothetical protein'; Neuros	36.5	96.6	54.4	154.8	1e-130	99 UNCLASSIFIED PROTEINS
Pc12g14890	strong similarity to fluconazole resistance protein f1e-123	0	T41018 probable membrane transporter - fission yeast (Schizosaccharomyces	12.0	66.9	17.8	95.2	1e-58	01.05.07 C-compound, carbohydrate transport
Pc12g16010	hypothetical protein	0	0	18.6	68.5	36.2	56.3	0	
Pc13g01890	strong similarity to acyl-CoA synthase like protein. 1e-103	0	B83121 probable AMP-binding enzyme PA4198 [imported] - Pseudomonas aer	145.0	223.6	132.0	391.1	1e-115	01 METABOLISM
Pc13g04680	strong similarity to alpha-cephem-methoxylase s1e-45	0	AP005958_178 gene: 'bld4449'; Bradyrhizobium japonicum USDA 110 DNA, complete	93.5	190.3	178.5	249.5	0	
Pc13g04990	strong similarity to hypothetical protein contig31_p 3e-12	0	AK051230_1 Mus musculus 12 days embryo spinal ganglion cDNA, RIKEN full-length	43.5	87.6	77.4	102.6	2e-47	13.05 cell motility
Pc13g07240	similarity to p-nitrobenzyl esterase pnbA - Bacillus 5e-77	0	T41347 probable carboxylesterase-lipase family member - fission yeast (Schiz	16.0	64.0	12.0	70.5	1e-113	13 REGULATION OF / INTERACTION WITH CELLULAR ENVIRONMENT
Pc13g07540	similarity to subunit 13 of glutathione S-transferase 1e-52	0	NCB7K22_4 gene: 'B7K22.040'; product: 'related to glutathione s-transferase subu	105.4	281.5	148.5	726.9	1e-87	01 METABOLISM
Pc13g08560	strong similarity to hypothetical protein An14g028c	0	0	12.1	38.5	31.8	20.4	0	
Pc13g10520	strong similarity to hypothetical protein yrmM - Bac 2e-82	0	BX842653_296 product: 'conserved hypothetical protein'; Belletobrio bacteriovorus c	42.7	74.9	43.2	87.1	1e-159	99 UNCLASSIFIED PROTEINS
Pc13g10810	strong similarity to acetoacetyl-CoA synthase acs3 0.0	0	BX649605_63 product: 'acetoacetyl-coa synthetase, putative'; Aspergillus fumigatus	108.2	224.2	213.9	409.9	1e-157	01 METABOLISM
Pc13g10880	hypothetical protein	0	0	12.0	133.9	14.2	37.0	0	
Pc13g10910	strong similarity to hypothetical protein An08g040f	0	0	56.3	170.2	82.4	126.8	0	
Pc13g11150	similarity to protein PRIB - Lentinus edodes 2e-11	0	PRIB_LENED PRIB PROTEIN.	12.7	81.1	29.5	67.7	0	
Pc13g11540	strong similarity to hypothetical oxidoreductase of 1e-39	0	AY458641_40 product: 'oxidoreductase, short-chain dehydrogenase/reductase family	12.0	80.7	13.2	12.8	6e-58	01 METABOLISM
Pc13g11580	strong similarity to hypothetical protein contig46_p 6e-87	0	BX649606_31 product: 'epoxide hydrolase, putative'; Aspergillus fumigatus BAC pilot	120.4	326.9	112.0	458.3	4e-66	01 METABOLISM
Pc13g11590	similarity to hypothetical protein SPCC1393.04 1e-10	0	T40952 hypothetical protein SPCC1393.04 - fission yeast (Schizosaccharomyc	48.5	133.4	39.5	98.1	0	
Pc13g11860	strong similarity to androgen-inducible aldehyde re 1e-166	0	BX649606_86 product: 'tafataxin b1 aldehyde reductase-like protein, putative'; Asper	131.9	282.7	255.7	271.0	4e-82	01 METABOLISM
Pc13g12110	strong similarity to hypothetical protein AAQ4946f 0.0	0	BX649605_68 product: 'hypothetical protein'; Aspergillus fumigatus BAC pilot project	75.1	153.5	134.6	175.8	0	
Pc13g13030	strong similarity to nonsense mRNA-reducing prot 9e-24	0	AE014829_55 product: 'regulator of nonsense transcripts, putative'; Plasmodium falci	12.0	17.1	12.0	25.3	1e-108	03.03.01 mitotic cell cycle and cell cycle control
Pc13g15930	similarity to hypothetical transcription regulator SP 6e-09	0	ENU56097_1 gene: 'fab3'; product: 'acetate regulatory DNA binding protein	34.4	149.9	22.8	165.3	1e-114	01.01.04 regulation of amino acid metabolism
Pc13g15940	strong similarity to copper amine oxidase AO1 - As 0.0	0	AF362473_1 product: 'copper amine oxidase'; Aspergillus niger strain AKU 3302 co	200.0	365.7	98.4	195.5	0.0	01 METABOLISM
Pc13g15950	strong similarity to choline permease Hnm1 - Sacc 1e-59	0	S11175 choline transport protein - yeast (Saccharomyces cerevisiae)	12.0	62.2	12.0	34.1	4e-78	01.01.07 amino acid transport
Pc16g00500	strong similarity to hypothetical membrane protein 1e-43	0	NC64C2_19 gene: '64C2.200'; product: 'related to PUTATIVE TARTRATE TRANS	33.5	70.5	178.1	716.8	0.0	08 CELLULAR TRANSPORT AND TRANSPORT MECHANISMS
Pc16g00610	strong similarity to hypothetical membrane protein 4e-91	0	CNS07TIX_2 DNA centromeric region sequence from BAC DP15B03, DP38F06 of cl	55.3	122.1	220.2	656.8	2e-71	01.01.07 amino acid transport
Pc16g00790	similarity to hypothetical transcription activator SP 2e-45	0	BX842620_25 gene: 'B11E5.250'; product: 'conserved hypothetical protein'; Neuros	35.2	89.0	53.8	98.2	1e-86	01.01.04 regulation of amino acid metabolism
Pc16g00830	similarity to hypothetical protein contig_1_108_scc	0	0	12.0	62.1	27.3	13.7	0	
Pc16g00840	similarity to O-methyltransferase A omtA - Aspergil 2e-13	0	BX908810_7 gene: 'G3C5.070'; product: 'related to O-methyltransferase'; Neurosp	35.3	121.2	83.0	86.1	3e-46	01.20 secondary metabolism
Pc16g00860	similarity to hypothetical protein contig17.fta_50cg	0	0	58.4	186.0	118.5	164.4	0	
Pc16g02950	strong similarity to aryl-alcohol oxidase precursor 4e-76	0	CNS07TIX_17 DNA centromeric region sequence from BAC DP15B03, DP38F06 of cl	19.2	22.1	18.7	45.5	0.0	02.11 electron transport and membrane-associated energy conservation
Pc16g04300	hypothetical protein	0	0	285.2	754.2	547.6	809.8	0	
Pc16g04830	strong similarity to pyridoxal 4-dehydrogenase Pl 6e-70	0	T39218 pyridoxal reductase - fission yeast (Schizosaccharomyces pombe)	12.0	24.4	24.6	34.8	4e-82	01.05.01.01.01 sugar, glucoside, polyol and carboxylate catabolism
Pc16g05120	weak similarity to hypothetical protein attM - Agrot 4e-08	0	AY052389_4 gene: 'attM/aiib'; product: 'AttM/Aiib'; Agrobacterium tumefaciens AttU	165.6	187.7	138.6	276.5	0	
Pc16g05680	strong similarity to alpha-L-arabinofuranosidase al 2e-95	0	G72395 alpha-L-arabinofuranosidase - Thermotoga maritima (strain MSB8)	112.9	128.1	38.3	198.5	0.0	01.05.01.01 C-compound, carbohydrate catabolism
Pc16g06760	strong similarity to hypothetical protein SC5H1.34 1e-45	0	T35350 hypothetical protein SC5H1.34 - Streptomyces coelicolor	27.4	115.5	57.9	124.7	5e-33	01.02 nitrogen and sulfur metabolism
Pc16g06970	strong similarity to short-chain alcohol dehydrogen 1e-28	0	APU76621_1 gene: 'adhA'; product: 'short-chain alcohol dehydrogenase'; Aspergill	21.8	27.5	23.7	60.2	3e-58	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc16g08590	similarity to NAD+-dependent 15-hydroxyprostagla 4e-22	0	CNS09S4S_94 DNA centromeric region sequence from BAC DP26B06, DP34F04, DP	51.3	81.4	33.7	75.8	1e-65	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc16g09230	strong similarity to urease URE1 - Cryptococcus n 0.0	0	U81509_1 gene: 'URE'; product: 'urease'; Coccidioides immitis urease (URE) ge	39.0	188.6	47.9	193.4	0.0	01.01.01.03.01 urea cycle
Pc16g10090	strong similarity to cytosine deaminase FCA1 - Ca 1e-46	0	AJ616007_1 gene: 'fca1'; product: 'putative cytosine deaminase'; Candida albican	75.2	152.8	154.6	294.6	1e-31	01.02 nitrogen and sulfur metabolism
Pc16g10640	weak similarity to transcription factor Arg81 - Sacc 8e-08	0	S49627 regulatory protein ARG82 - yeast (Saccharomyces cerevisiae)	26.2	64.0	54.8	131.0	0.0	01.01.04 regulation of amino acid metabolism
Pc16g11760	weak similarity to hypothetical transcription regula 2e-04	0	T37601 probable transcription factor - fission yeast (Schizosaccharomyces pom	92.4	197.9	187.7	217.1	2e-60	01.01.04 regulation of amino acid metabolism
Pc16g11770	strong similarity to choline monoxygenase like pn 3e-27	0	BX248345_40 gene: 'Mb3186c'; product: 'POSSIBLE DIOXYGENASE'; Mycobacter	45.0	125.6	56.9	89.8	1e-148	01.20.17.03 biosynthesis of amines
Pc16g12280	strong similarity to membrane protein Tpo2 - Sacc 0.0	0	BX649607_11 gene: 'mdr1'; product: 'mfs-family multidrug resistance protein, putative	37.5	291.2	103.4	364.0	2e-63	01.05.07 C-compound, carbohydrate transport
Pc16g12810	strong similarity to hypothetical protein CAC2874a 6e-76	0	BX649607_60 product: 'Hypothetical protein, conserved'; Aspergillus fumigatus BAC	561.7	691.9	537.0	1168.0	0	
Pc16g13180	strong similarity to L-galactonolactone oxidase Aio 2e-90	0	AF031228_1 gene: 'ALO'; product: 'D-arabinono-1,4-lactone oxidase'; Candida alb	35.1	99.5	58.8	92.0	1e-171	01.05.01 C-compound and carbohydrate utilization
Pc16g13370	similarity to hypothetical protein contig_1_13_scaf 2e-04	0	AC115684_23 product: 'hypothetical protein'; Dictyostelium discoideum chromosome	23.7	70.3	42.4	71.4	0	
Pc16g13490	strong similarity to short branched chain specific a 1e-121	0	HSM803134_1 gene: 'DKFZp451K202'; product: 'hypothetical protein'; Homo sapien	301.5	432.9	290.4	744.8	1e-138	01.01.10.04 degradation of amino acids of the pyruvate family
Pc16g13540	similarity to positive regulator of purine utilisation 4e-30	0	SA5779 purine utilization positive regulator - Emericella nidulans	12.0	39.3	12.0	46.8	1e-100	01.01.04 regulation of amino acid metabolism
Pc17g00430	weak similarity to hypothetical membrane protein 1	0	0	184.8	562.3	238.2	264.8	0	
Pc17g00440	weak similarity to retrotransposon Tto1 - Nicotian 1e-145	0	T02206 hypothetical protein - common tobacco retrotransposon Tto1	38.7	309.2	32.6	113.1	0.0	10.01.01 unspecified signal transduction
Pc17g00590	strong similarity to retrotransposon Tto1 - Nicotian 1e-140	0	T02206 hypothetical protein - common tobacco retrotransposon Tto1	65.5	470.5	45.4	153.2	0.0	10.01.01 unspecified signal transduction
Pc17g00590	strong similarity to retrotransposon Tto1 - Nicotian 1e-140	0	T02206 hypothetical protein - common tobacco retrotransposon Tto1	65.5	470.5	45.4	153.2	0.0	10.01.01 unspecified signal transduction
Pc18g00150	strong similarity to high affinity glucose transporte 3e-38	0	AF136235_1 gene: 'asd-3'; product: 'ascus development protein 3'; Neurospora cr	39.1	113.2	117.6	68.9	9e-76	01.05.07 C-compound, carbohydrate transport
Pc18g00160	strong similarity to hypothetical protein 1357_scaf	0	0	37.9	92.3	62.7	62.3	0	
Pc18g00500	strong similarity to hypothetical protein contig336.14e-99	0	BX842620_43 gene: 'B11E5.430'; product: 'related to dihydropicolinate synthase';	70.5	114.4	68.1	149.5	9e-92	01.01.01.07.06 biosynthesis of lysine
Pc18g02660	strong similarity to hypothetical protein An18g040f	0	0	17.6	39.5	21.4	17.2	0	
Pc18g02670	strong similarity to salicylaldehyde dehydrogenase 5e-96	0	ACPCAOP_34 gene: 'hcaB'; product: 'vanillin dehydrogenase'; Acinetobacter sp. ADI	143.3	374.7	102.8	332.4	0.0	01.05 C-compound and carbohydrate metabolism
Pc18g03010	strong similarity to choline permease Hnm1 - Sacc 1e-75	0	S11175 choline transport protein - yeast (Saccharomyces cerevisiae)	60.3	125.0	36.8	75.2	2e-75	01.01 amino acid metabolism
Pc18g05670	strong similarity to hypothetical protein SPCC16A12e-23	0	T41080 hypothetical protein SPCC16A11.07 - fission yeast (Schizosaccharomyc	58.1	92.4	51.6	113.1	7e-44	04.05.01.04 transcriptional control

Pc18g05780	strong similarity to multidrug resistance protein fnx9e-79	T40380	major facilitator family transporter - fission yeast (Schizosaccharomyces pombe)	101.4	283.6	218.6	530.3	1e-49	01.05.04 regulation of C-compound and carbohydrate utilization
Pc18g06290	strong similarity to hypothetical NADPH dehydrog 1e-104	BX908808_9	gene: "G2B184.090"; product: "probable NADPH2 dehydrogenase chain 2"	12.1	45.6	26.0	65.1	3e-81	01.20 secondary metabolism
Pc19g00090	weak similarity to retrotransposon Tto1 - Nicotiana glauca	T02206	hypothetical protein - common tobacco retrotransposon Tto1	13.6	29.3	13.1	16.2	0.0	10.01.01 unspecified signal transduction
Pc19g00550	strong similarity to cutinase transcription factor bel1e-56	CT1B_FUSSO	CUTINASE TRANSCRIPTION FACTOR 1 BETA.	226.3	450.3	165.4	332.6	9e-68	01.01.04 regulation of amino acid metabolism
Pc20g00130	strong similarity to myo-inositol transport protein h1e-141	NCB17B1_7	gene: "B17B1.070"; product: "related to myo-inositol transporter"; Neurospora crassa	168.2	567.6	149.7	679.4	2e-72	01.05.07 C-compound, carbohydrate transport
Pc20g01500	strong similarity to 2,3-dihydroxybenzoic acid decarboxylase 1e-69	AF2887	conserved hypothetical protein ALU2529 [imported] - Agrobacterium tumefaciens	76.3	292.0	224.7	280.1	1e-133	99 UNCLASSIFIED PROTEINS
Pc20g01980	strong similarity to phenol 2-monooxygenase - Trifolium pratense	AY450844_1	product: "n-hydroxybenzoate hydroxylase"; Comamonas testosteroni	12.0	60.6	12.4	79.5	9e-92	01.01.10.05.07 degradation of tryptophan
Pc20g02630	strong similarity to Gly-X carboxypeptidase precun5e-85	SCCP51_1	gene: "CP51"; product: "carboxypeptidase S"; Saccharomyces cerevisiae	17.8	38.4	21.0	104.5	1e-48	01.01 amino acid metabolism
Pc20g04510	strong similarity to multidrug resistance protein Hoi1 - Saccharomyces cerevisiae	T39346	probable major facilitator family multi-drug resistance protein - fission yeast (Schizosaccharomyces pombe)	12.0	43.5	12.0	41.7	3e-47	01.05.04 regulation of C-compound and carbohydrate utilization
Pc20g06080	strong similarity to hypothetical protein An02g114k.0.0	NCB19A17_11	gene: "B19A17.110"; product: "hypothetical protein"; Neurospora crassa	203.1	271.4	111.2	630.0	5e-86	01.03 nucleotide metabolism
Pc20g06140	strong similarity to hypothetical protein An02g113z.1e-108	NCB19A17_12	gene: "B19A17.120"; product: "hypothetical protein"; Neurospora crassa	203.1	271.4	111.2	630.0	5e-86	01.03 nucleotide metabolism
Pc20g06200	strong similarity to hypothetical membrane protein 1e-118	S50361	probable membrane protein YIL166c - yeast (Saccharomyces cerevisiae)	12.0	25.6	95.6	525.6	2e-99	01.01.04 regulation of amino acid metabolism
Pc20g06650	hypothetical protein	0	0	136.4	162.0	113.6	281.5	0	
Pc20g07520	similarity to hypothetical protein An01g15120 - Aspergillus fumigatus	BX649605_78	product: "hypothetical protein, conserved"; Aspergillus fumigatus BAC	127.2	105.9	88.0	242.1	0	
Pc20g08300	strong similarity to hypothetical protein 12F11.2005e-59	NC12F11_20	gene: "12F11.200"; product: "hypothetical protein"; Neurospora crassa	51.2	76.8	72.3	150.1	0	
Pc20g08400	similarity to hypothetical transcriptional regulator F2e-12	AF057038_1	gene: "FCR1"; product: "zinc cluster transcription factor Fcr1p"; Candida albicans	26.4	63.4	58.0	95.1	6e-19	01.01.04 regulation of amino acid metabolism
Pc20g08470	strong similarity to protease Mch5 - Saccharomyces cerevisiae	NCB23B10_3	gene: "B23B10.030"; product: "related to monocarboxylate transporter"	21.2	64.9	77.5	239.3	1e-112	01 METABOLISM
Pc20g10520	similarity to hypothetical protein An12g07270 - Aspergillus fumigatus	0	0	12.0	22.2	13.7	29.6	0	
Pc20g12910	strong similarity to hypothetical protein An08g121z	0	0	25.2	63.0	45.6	34.4	0	
Pc20g14130	strong similarity to hypothetical protein 1160_scaff1e-156	AP004600_12	gene: "OB2083"; product: "hypothetical conserved protein"; Oceanobacillus anophagealis	111.0	124.2	84.3	186.4	1e-68	01.05 C-compound and carbohydrate metabolism
Pc20g14390	strong similarity to mitochondrial phosphate transp 1e-97	S50556	hypothetical protein YER053c - yeast (Saccharomyces cerevisiae)	12.0	65.9	35.6	233.5	1e-87	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc20g16550	strong similarity to hypothetical conserved protein 1e-22	E82998	conserved hypothetical protein PA5185 [imported] - Pseudomonas aer	97.4	132.5	117.5	237.4	0	
Pc21g02160	weak similarity to TRAF5 - Mus musculus	NCB23E9_9	gene: "B23E9.090"; product: "related to traf5 protein"; Neurospora cr	83.7	159.1	81.4	177.5	6e-81	04.05.01.04 transcriptional control
Pc21g03190	strong similarity to hydroxy-pyruvate dehydrogenase 6e-72	BX649606_114	product: "glycerate dehydrogenase, putative"; Aspergillus fumigatus B	93.1	111.2	72.9	189.5	1e-100	01 METABOLISM
Pc21g04410	strong similarity to glucose 1-dehydrogenase ghdl 9e-47	TACID3_96	gene: "Ta0747"; product: "glucose 1-dehydrogenase related protein";	12.0	13.7	12.0	27.8	6e-63	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc21g05110	strong similarity to hypothetical protein contig_1.54e-79	T49756	related to berberine bridge enzyme [imported] - Neurospora crassa	27.0	207.9	151.1	66.5	1e-100	01.05.01 C-compound and carbohydrate utilization
Pc21g05550	strong similarity to multidrug resistance protein fnx5e-70	NCB1308_14	gene: "B1308.150"; product: "conserved hypothetical protein"; Neuros	42.6	156.1	45.0	104.4	5e-50	01.05.04 regulation of C-compound and carbohydrate utilization
Pc21g05750	strong similarity to fatty acid regulation protein lke8e-49	NCB9B15_9	gene: "BB15.090"; product: "conserved hypothetical protein"; Neuros	901.2	2504.4	1322.7	3172.5	2e-76	01.06.10 regulation of lipid, fatty-acid and isoprenoid metabolism
Pc21g05850	strong similarity to allantoinase permease Dals - Saccharomyces cerevisiae	T41604	probable membrane transport protein - fission yeast (Schizosaccharom	21.9	100.6	127.3	554.3	1e-104	08 CELLULAR TRANSPORT AND TRANSPORT MECHANISMS
Pc21g06300	strong similarity to hypothetical protein contg5_p3e-84	BX908808_19	gene: "G2B184.190"; product: "related to NADPH2 quinone reductase";	111.9	515.6	233.5	1088.0	9e-74	01 METABOLISM
Pc21g06670	strong similarity to carboxypeptidase S1 - Penicillium chrysogenum	AF394242_1	gene: "cpl"; product: "carboxypeptidase S1"; Aspergillus oryzae strain 1	53.3	52.0	30.5	170.3	1e-105	06.07 protein modification
Pc21g07740	similarity to beta-ketoadipyl reductase rhG5 - Pseudomonas fluorescens	NCB23B10_7	gene: "B23B10.070"; product: "related to gluconate 5-dehydrogenase";	143.5	317.3	240.3	541.0	8e-67	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc21g07900	strong similarity to hypothetical protein contig_1.1	0	0	295.9	504.4	292.6	594.1	0	
Pc21g08400	strong similarity to 3-hydroxyisobutyryl-coenzyme 1e-137	NCB18D24_21	gene: "B18D24.230"; product: "related to enoyl-CoA hydratase"; Neuro	92.2	206.2	88.2	379.8	1e-111	01 METABOLISM
Pc21g08920	strong similarity to norsolorinic acid reductase nor1e-119	AP24698_1	gene: "nor"; product: "norsolorinic acid reductase"; Aspergillus parasit	78.1	294.0	333.2	414.8	8e-88	01.01.10 amino acid degradation (catabolism)
Pc21g10100	weak similarity to glutathione S-transferase omeg3e-11	TCTCAC2X_1	product: "TcAc2"; Trypanosoma cruzi stress and GST superfamily rela	131.1	637.8	260.2	1015.8	4e-41	03.01.05.01 DNA repair
Pc21g12380	strong similarity to hypothetical neutral amino acid 2e-66	S47892	neutral amino acid permease - Neurospora crassa	33.2	74.7	56.8	133.9	2e-67	01.01.07 amino acid transport
Pc21g13520	weak similarity to hypothetical protein YKL051w - Aspergillus fumigatus	NCB2F7_3	gene: "B2F7.030"; product: "conserved hypothetical protein"; Neurosp	123.3	272.4	136.5	167.6	0	
Pc21g14050	strong similarity to mannase man1 - Aspergillus aculeatus	AAMANNA_1	gene: "man1"; product: "mannanase"; Aspergillus aculeatus mannans	56.7	48.2	35.7	77.0	1e-123	01.05 C-compound and carbohydrate metabolism
Pc21g14230	strong similarity to hypothetical nadh-dependent fl 5e-89	T39956	probable nadh-dependent flavin oxidoreductase - fission yeast (Schiz	74.5	266.9	249.7	775.7	0	
Pc21g14260	strong similarity to high-affinity nicotinic acid perm 1e-120	SPBC1683_12	gene: "SPBC1683.12"; Schizosaccharomyces pombe cosmid c1683.	43.6	88.4	407.3	1042.9	1e-84	01.01.10 amino acid degradation (catabolism)
Pc21g14270	similarity to hypothetical transcription regulator SP2e-16	T40521	hypothetical protein SPBC530.05 - fission yeast (Schizosaccharomyces	67.2	256.1	91.8	245.5	8e-91	01.01.04 regulation of amino acid metabolism
Pc21g14290	strong similarity to hypothetical protein An12g073z	0	0	12.0	23.2	12.0	36.6	0	
Pc21g14380	similarity to hypothetical protein SPBC31F10.02 - Aspergillus fumigatus	NC18F11_2	gene: "18F11.015"; product: "conserved hypothetical protein"; Neuros	168.2	370.3	196.8	370.7	8e-40	47.03.01.01.01.99 other brain areas
Pc21g14740	strong similarity to hypothetical intracellular protease am3e-17	G87469	Thl1/Ptp1 family protein [imported] - Caulobacter crescentus	30.6	82.0	35.4	96.9	0	
Pc21g16710	weak similarity to hypothetical protein DR0703 - Drosophila melanogaster	H69851	hypothetical protein yJlB - Bacillus subtilis	22.2	72.7	14.7	63.0	3e-53	99 UNCLASSIFIED PROTEINS
Pc21g17040	weak similarity to beta transducin-like protein het-5e-05	AF323585_1	gene: "het-d"; product: "beta transducin-like protein HET-D2Y"; Podo	89.4	176.1	68.5	409.3	0	
Pc21g19280	strong similarity to 25D9-5p - Aspergillus fumigatus	AY080962_7	gene: "25d9-5"; product: "25D9-5p"; Aspergillus fumigatus Mvp1p (Zod	43.9	139.3	27.8	160.0	0	
Pc21g20580	strong similarity to Fe(II)-dependent sulfonate alpha 7e-55	S50963	hypothetical protein YLL057c - yeast (Saccharomyces cerevisiae)	164.1	192.9	142.1	717.9	1e-152	11.07.01 detoxification involving cytochrome P450
Pc21g20650	strong similarity to 4-coumarate-CoA ligase 4CL - Escherichia coli	AP005036_223	product: "putative 4-coumarate:CoA ligase"; Streptomyces avermectin	72.6	62.9	52.6	123.1	1e-125	01 METABOLISM
Pc21g20890	strong similarity to NADPH cytochrome P450 oxidoreductase 0.0	AX684154_1	Sequence 5 from Patent WO0246386.	912.0	2551.3	626.1	2715.6	0.0	01.02.01.07 nitric oxide biosynthesis
Pc21g21340	similarity to hypothetical protein alr1138 - Nostoc sp. (strain PCC 7120)	AG1948	hypothetical protein alr1138 [imported] - Nostoc sp. (strain PCC 7120)	12.1	51.8	12.0	23.1	0	
Pc21g21350	weak similarity to hypothetical conserved protein F2e-28	NC5F3_10	gene: "5F3.100"; product: "conserved hypothetical protein"; Neurosp	127.1	741.5	585.0	1037.2	0	
Pc21g23440	strong similarity to hypothetical aminotransferase 1e-144	BX897674_17	gene: "B23B18.180"; product: "conserved hypothetical protein"; Neuros	201.0	388.1	66.6	213.0	1e-133	01.01 amino acid metabolism
Pc21g23730	strong similarity to feruloyl-CoA synthase like proti 0.0	CNS07ITX_8	DNA centromeric region sequence from BAC DP15B03, DP38F06 of chromo	36.7	81.0	52.3	129.8	1e-143	01 METABOLISM
Pc22g00060	similarity to fructosyl amine oxygen oxidoreductase 1e-174	AB035128_1	gene: "sox"; product: "sarcosine oxidase"; Cylinodrocarpon didymus s	93.9	177.6	137.9	334.2	1e-100	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc22g00110	strong similarity to hypothetical oxidoreductase SF1e-105	AE119281_1	Alternaria alternata AMT2 gene for aldo-keto reductase, complete	100.5	99.2	28.9	532.6	5e-87	01.01.10 amino acid degradation (catabolism)
Pc22g00130	strong similarity to hypothetical protein 1110_scaff1e-50	BX842620_43	gene: "B11E5.430"; product: "related to dihydroadipic acid synthase";	25.4	96.0	34.0	125.4	4e-83	01.01.01.07.06 biosynthesis of lysine
Pc22g00140	strong similarity to exo-alpha-sialidase - Micromonospora viridifaciens	A45244	exo-alpha-sialidase (EC 3.2.1.18) - Micromonospora viridifaciens	12.9	72.6	16.6	74.4	1e-137	01.05 C-compound and carbohydrate metabolism
Pc22g00270	strong similarity to transcription activator Put3 - Saccharomyces cerevisiae	A39792	transcription activator PUT3 - yeast (Saccharomyces cerevisiae)	99.3	208.8	140.7	337.3	1e-92	01.01.04 regulation of amino acid metabolism
Pc22g01120	strong similarity to hypothetical protein PA2326 - Pseudomonas aeruginosa	A83356	hypothetical protein PA2326 [imported] - Pseudomonas aeruginosa (sti	71.3	131.8	135.5	830.8	3e-51	01.05.07 C-compound, carbohydrate transport
Pc22g01340	strong similarity to D-arabinol 2-dehydrogenase 1e-37	D70635	hypothetical protein Rv1928c - Mycobacterium tuberculosis (strain H37	12.0	31.5	13.3	28.6	3e-64	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc22g01530	strong similarity to hypothetical protein YAL049c - Aspergillus fumigatus	AY465528_1	product: "diene lactone hydrolase"; Chaetomium globosum diene lacton	234.0	787.2	740.1	845.0	2e-68	98 CLASSIFICATION NOT YET CLEAR-CUT
Pc22g01540	strong similarity to hypothetical protein SPBC3B8 - Schizosaccharomyces pombe	CNS09S45_72	DNA centromeric region sequence from BAC DP26B06, DP34F04, DP	447.3	654.3	176.2	454.6	8e-90	02.13 respiration
Pc22g01660	similarity to uroporphyrinogen-III synthase HemA - Escherichia coli	CNS09S45_38	DNA centromeric region sequence from BAC DP26B06, DP34F04, DP	30.9	32.3	17.0	49.6	4e-92	01.07.01 biosynthesis of vitamins, cofactors, and prosthetic groups
Pc22g02620	strong similarity to monoglyceride lipase mgl1 - Mus musculus	NC7F4_20	gene: "7F4.210"; product: "conserved hypothetical protein"; Neurosp	71.7	119.1	89.4	238.3	9e-41	01.05.01.01.09 aerobic aromatic catabolism
Pc22g02720	weak similarity to hypothetical conserved protein 1e-61	NC104H10_16	gene: "104H10.140"; product: "hypothetical protein"; Neurospora cras	29.9	26.7	21.9	45.9	0	
Pc22g030710	strong similarity to hypothetical membrane protein 1e-168	NCB2P8_18	gene: "BP8.200"; product: "related to dihydroxy transporter"; Neuros	14.3	49.7	26.5	82.4	3e-59	01.05.07 C-compound, carbohydrate transport
Pc22g03940	strong similarity to Succinyl-CoA:3-oxoacid CoA transferase 1e-155	AB103529_1	product: "Homo sapiens 3-oxoacid CoA transferase"; Synthetic constr	204.4	697.9	170.5	775.0	0.0	01 METABOLISM
Pc22g04190	similarity to protein Mop1 - Saccharomyces cerevisiae	SCF0993	hypothetical protein YJR074w - yeast (Saccharomyces cerevisiae)	35.4	46.6	36.6	75.5	5e-13	06.04 protein targeting, sorting and translocation
Pc22g05400	strong similarity to hypothetical monocarboxylate transporter 1e-46	S38065	hypothetical protein YKL221w - yeast (Saccharomyces cerevisiae)	22.7	89.1	67.2	266.8	0	
Pc22g06010	similarity to hypothetical protein YBR151w - Saccharomyces cerevisiae	S46022	hypothetical protein YBR151w - yeast (Saccharomyces cerevisiae)	39.9	74.6	42.9	96.4	7e-15	01.05 C-compound and carbohydrate metabolism
Pc22g06830	similarity to inhibitor of endosome-lysosome fusion 4e-16	AP005027_159	product: "putative dimethylmenaquinone methyltransferase family prote	33.2	56.6	22.2	90.6	6e-81	99 UNCLASSIFIED PROTEINS
Pc22g07330	strong similarity to hypothetical protein An16g068z 2e-09	AF230811_1	gene: "PIG1"; product: "putative transcription factor Pig1p"; Pityriaria	131.8	309.5	103.9	296.7	6e-96	04.05.01.04 transcriptional control
Pc22g07440	strong similarity to benzoate 4-monooxygenase cy 1e-154	NCB14A6_3	gene: "B14A6.030"; product: "probable benzoate 4-monooxygenase cy	25.9	76.4	57.9	88.0	1e-101	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc22g09840	strong similarity to cytochrome P450 monooxygenase 4e-92	T49413	prob						

Pc22g15710	similarity to hypothetical protein An08g08410 - Asq	0	0	0	24.4	66.0	62.6	91.6	0	
Pc22g16060	strong similarity to UDP-glucose:dolichyl-phosphat	T44819	dolichyl-phosphate beta-glucosyltransferase (EC 2.4.1.117) [imported]	92.4	169.3	84.2	287.0	2e-85	01 METABOLISM	0
Pc22g17060	strong similarity to hypothetical protein An12g1040	0	0	46.3	50.9	27.5	59.7	0		0
Pc22g17250	strong similarity to hypothetical protein mg02084.12e-32	D86873	dehydrogenase [imported] - Lactococcus lactis subsp. lactis (strain IL1	85.3	163.0	131.0	277.1	1e-104	01 METABOLISM	0
Pc22g17810	weak similarity to hypothetical protein yciI - Eschei 4e-19	NC20H10_30	gene: "20H10.300"; product: "conserved hypothetical protein"; Neuros	280.3	840.9	455.9	1365.2	0		0
Pc22g18380	strong similarity to salicylate hydroxylase nahG - F2e-35	AP005214_117	product: "putative salicylate hydroxylase"; Corynebacterium efficiens Y	14.5	40.3	38.9	68.7	2e-85	01.01.10 amino acid degradation (catabolism)	0
Pc22g19410	weak similarity to hypothetical protein YJR154w - 4e-89	SCO939129_22	gene: "SCO6985"; "SC8F11.11c"; product: "hypothetical protein SC8F	15.1	20.6	12.0	29.0	0		0
Pc22g20490	strong similarity to cinnamyl-alcohol dehydrogenat	NCB23E9_11	gene: "B23E9.110"; product: "conserved hypothetical protein"; Neuros	158.3	368.5	291.4	357.4	5e-52	01.03 nucleotide metabolism	0
Pc22g21310	weak similarity to hypothetical protein 192_scaffol	0	0	15.5	149.1	113.6	269.2	0		0
Pc22g22590	similarity to alcohol dehydrogenase pan2 - Mus m 8e-18	BC063926_1	gene: "MGC76232"; product: "MGC76232 protein"; Xenopus tropicalis	17.3	71.7	54.2	44.7	3e-56	01 METABOLISM	0
Pc22g24570	similarity to putative integral membrane protein C7 5e-71	BX908810_10	gene: "G3C5.100"; product: "related to integral membrane protein PTH	16.6	16.3	12.6	48.0	0		0
Pc22g24790	similarity to cutinase transcription factor alpha CTI3e-30	T50198	probable transcription activator protein [imported] - fission yeast (Schiz	51.1	120.9	73.1	91.0	1e-126	01.01.04 regulation of amino acid metabolism	0
Pc22g25120	strong similarity to hypothetical protein 1141_scaff 7e-67	BX572594_63	product: "probable enoyl CoA-hydratase/isomerase"; Rhodospseudomc	12.0	20.1	14.8	38.2	1e-71	01 METABOLISM	0
Pc22g26000	weak similarity to retrotransposon Tto1 - Nicotians 7e-61	S04273	retrovirus-related reverse transcriptase homolog - common tobacco ret	19.3	59.6	33.8	38.2	0.0	10.01.01 unspecified signal transduction	0
Pc22g26000	weak similarity to retrotransposon Tto1 - Nicotians 7e-61	S04273	retrovirus-related reverse transcriptase homolog - common tobacco ret	19.3	59.6	33.8	38.2	0.0	10.01.01 unspecified signal transduction	0
Pc24g03100	weak similarity to retrotransposon Tto1 - Nicotians 1e-137	T02206	hypothetical protein - common tobacco retrotransposon Tto1	89.6	230.0	83.8	95.1	0.0	10.01.01 unspecified signal transduction	0

Supplementary Table 11. K-mean cluster 3
@=Values given are the average of three independent experiments

Ref code	Description of putative <i>P. chrysogenum</i> ORF	Best blast homolog to putative <i>P. chrysogenum</i> ORF	Gene	description	WS - PAA	DS - PAA	LS - PAA	DS + PAA	Prnt cat (rate)	Category
P060360	hypothetical protein	7e-04	AB068070	product: "envelope glycoprotein precursor". Cme16n-Congo he	12.0	20.8	72.9	123.0		
P060361	strong similarity to high affinity glucose transporter HGT1 - Kluyveromyces fragilis	6e-59	AY081846	gene: "m8B"; product: "putative sugar transporter"; Aspergillus	12.0	7.3	12.0	12.0	26-69	01.05.04 regulation of C-compound and carbohydrate utilization
P060362	strong similarity to maltose transport protein Mal1 - Saccharomyces cerevisiae	1e-86	NC55E.16	gene: "SE6.100"; product: "probable alpha-glucoside transport i	27.9	9.14	45.8	161.1	36-82	01.05.07 C-compound, carbohydrate transport
P060363	strong similarity to lysine permease YAP1 - Saccharomyces cerevisiae	1e-123	AY072168	gene: "YAP1"; product: "unknown"; Aspergillus fumigatus	12.0	14.0	12.0	12.0	11-4	01.05.04 regulation of C-compound and carbohydrate utilization
P060364	strong similarity to quinone to quinoxaline reductase P1 - Arabidopsis thaliana	2e-82	AF525099	gene: "red1"; product: "reductase RED1"; Cochliobolus hetero	81.9	9.3	142.0	586.4	79-9	01.01.01 amino acid metabolism
P060365	strong similarity to oxalacetate to short-chain ORF1 - Streptomyces coelicolor	1e-47	T41028	short chain dehydrogenase - fission yeast (Schizosaccharomyces	12.0	12.0	18.8	90.1	16-80	01.05.02 C-compound and carbohydrate metabolism
P060366	weak similarity to hypothetical protein PA2338 - Pseudomonas aeruginosa	7e-17	BX49807	product: "hypothetical protein"; Aspergillus fumigatus	12.0	12.0	12.0	12.0	12-1	
P060367	strong similarity to multistep resistance protein frx1p - Saccharomyces pombe	1e-178	NC55E.16	gene: "frx1B308.100"; product: "frx1B308.100"; Aspergillus	12.0	12.0	12.0	12.0	12-1	01.05.07 C-compound, carbohydrate transport
P060368	strong similarity to aldehyde reductase Hra - Sporobolus salmonicolor	2e-92	AF167899	gene: "HraR"; product: "aldehyde reductase Hra"; Sporobolus sal	153.2	66.0	170.6	1867.3	5-13	01.05.02 C-compound, carbohydrate metabolism
P060369	strong similarity to methylglyoxal resistance gene AMR1 - Staphylococcus aureus	1e-100	AF4241.9	gene: "B1A1.240"; product: "related to amyloid hydrolysis AnHx	12.0	12.0	12.0	12.0	12-1	01.05.02 C-compound and carbohydrate metabolism
P060370	strong similarity to valyl dihydroxyase Acs - Galactospora strobilata	2e-54	AF0289.9	product: "valyl dihydroxyase Acs"; Hypocrea conio	12.0	12.0	12.0	12.0	12-1	01.05.07 C-compound, carbohydrate transport
P060371	strong similarity to p-nitrobenzyl esterase pNB4 - Bacillus subtilis	1e-130	BS272606	gene: "estA1"; product: "type B carboxylesterase"; Bacillus s	12.0	12.0	12.0	12.0	12-1	01.04.07 phosphate metabolism
P060372	weak similarity to hypothetical isochromatase Ta0728 - Thermoplasma acidophilum	4e-24	F83436	conserved hypothetical protein PA1077 [imported] - Pseudomon	40.8	113.6	156.8	1249.6	26-47	01.01.11 biosynthesis of the cysteine-aromatic group
P060373	strong similarity to hypothetical protein PA1077 - Pseudomonas aeruginosa	4e-24	AB071807	product: "hypothetical protein"; Aspergillus fumigatus	12.0	12.0	12.0	12.0	12-1	01.01.09 amino acid metabolism activities
P060374	strong similarity to polyamine transport protein Pto1 - Saccharomyces cerevisiae	5e-22	BX49807	product: "H17A4.250"; product: "related to multistep resistant pro	19.4	21.0	36.6	102.3	76-45	01.04.07 phosphate transport
P060375	strong similarity to hypothetical protein PA1077 - Pseudomonas aeruginosa	5e-22	T41000	product: "SPAC222.050"; product: "fission yeast (Schizosacchar	12.0	12.0	12.0	12.0	12-1	01.03.03 nucleotide metabolism
P060376	strong similarity to extracellular alpha-glucosidase AgU1 - Aspergillus niger	1e-10	JC4217	alpha-glucosidase (EC 3.2.1.20) - Aspergillus oryzae	24.7	32.9	105.0	109.3	10-3	01.01.01 amino acid biosynthesis
P060377	strong similarity to hypothetical membrane protein YGL1 - Saccharomyces cerevisiae	2e-57	NC528310	gene: "YGL1.030"; product: "related to noncarboxylesterase trans	12.0	12.0	12.0	12.0	12-1	
P060378	strong similarity to hexameric surface lectin - Aspergillus fumigatus	3e-18	AF1769	product: "hexameric surface lectin - Aspergillus fumigatus (Af)	12.0	12.0	12.0	12.0	46-55	01.05.04 regulation of C-compound and carbohydrate utilization
P060379	strong similarity to integral membrane protein Mgt11 - Magnaporthe oryzae	3e-30	AF119671	gene: "PTPH1.1"; product: "integral membrane protein"; Magnap	90.1	757.9	1558.8	3036.3	26-83	01.07.01 biosynthesis of vitamins, cofactors, and prosthetic groups
P060380	strong similarity to hypothetical protein CC3654 - Caulobacter crescentus	2e-23	BX569693	product: "conserved hypothetical protein"; Synedochys sp. V	12.0	12.0	12.0	12.0	12-1	01.05.02 C-compound and carbohydrate metabolism
P060381	weak similarity to 2-hydroxyisovaleryl-CoA lyase BL2 - Bacillus subtilis	1e-120	AF49105	product: "2-hydroxyisovaleryl-CoA lyase BL2"; Bacillus subtilis	22.8	90.3	73.8	300.3	2-82	01.05.02 C-compound and carbohydrate metabolism
P060382	strong similarity to benzoylformate decarboxylase - Pseudomonas putida	1e-120	NC82014.1	gene: "BZD14.130"; product: "related to BENZOYLFORMATE D	12.0	12.0	12.0	12.0	12-1	01.05.02 C-compound and carbohydrate metabolism
P060383	strong similarity to isochrysoerythrin lyase PAX111 - Pseudomonas aer	3e-16	F83006	isochrysoerythrin lyase PAX111 [imported] - Pseudomonas aer	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060384	strong similarity to hypothetical protein PA1077 - Pseudomonas aeruginosa	1e-120	AB071807	product: "hypothetical protein"; Aspergillus fumigatus	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060385	strong similarity to hypothetical protein encoded by at3139 - Nostoc sp.	1e-23	D83426	hypothetical protein PA1752 [imported] - Pseudomonas aerugin	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060386	strong similarity to starvation-sensing protein like AnG042760 - Aspergillus niger	1e-112	AY040060	gene: "OB252.15"; product: "unknown conserved protein (partial)	12.0	12.0	12.0	12.0	12-1	01.01.07 amino acid transport
P060387	strong similarity to alcohol dehydrogenase adhP - Pseudomonas aeruginosa	1e-108	AY170473	product: "alcohol dehydrogenase adhP"; Pseudomonas aerugin	12.0	12.0	12.0	12.0	12-1	01.05.01.01.01 sugar, glucoside, polyol and carbohydrate catabolism
P060388	weak similarity to hypothetical oxoacidase PA3530 - Pseudomonas aeruginosa	2e-24	NC55F.10	gene: "SF3.100"; product: "conserved hypothetical protein"; Ne	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060389	strong similarity to quinate transport protein qut - Aspergillus nidulans	1e-61	S08498	quate transport protein - Emmericella nidulans	180.7	231.0	142.9	360.7	26-50	01.05.04 regulation of C-compound and carbohydrate utilization
P060390	strong similarity to urea transport protein Urt2 - Saccharomyces cerevisiae	1e-114	SA6330	urea transport protein - yeast (Saccharomyces cerevisiae)	84.7	99.3	126.0	126.0	12-1	01.04.07 phosphate transport
P060391	strong similarity to lysine permease YAP1 - Saccharomyces cerevisiae	1e-130	BX080808	product: "YAP1.030"; product: "probable amino acid transporter";	84.7	99.3	126.0	126.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060392	strong similarity to polyamine transport protein Pto1 - Saccharomyces cerevisiae	1e-22	AP005024	product: "hypothetical protein"; Streptomyces avermitilis genom	122.5	288.4	350.2	928.6	6-6	01.01.01.01.02 biosynthesis of the glutamate group (proline, hydroxyproline, arginine, glutamine, glutamate)
P060393	strong similarity to hypothetical conserved protein SPH2.80 - Neurospora crassa	1e-130	BX498025	product: "hypothetical protein, conserved"; Aspergillus fumigat	18.6	27.3	61.8	96.21	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060394	strong similarity to hypothetical protein Aka2 - Aspergillus fumigatus	3e-09	AY164464	gene: "Pto103"; product: "polyspecific ketone diase, f-ike 3 - M	13.7	24.1	71.2	110.2	96-7	99 UNCLASSIFIED PROTEINS
P060395	strong similarity to mycobiosyl transferase ynfB - Saccharomyces cerevisiae	3e-63	AE2201	hypothetical protein aif723 [imported] - Nostoc sp. (strain PCC	16.9	47.8	11.1	973.9	86-9	01.05.02 C-compound and carbohydrate metabolism
P060396	weak similarity to phenylacetylserine lyase - Aspergillus fumigatus	2e-78	BX498025	product: "hypothetical protein"; Aspergillus fumigatus	12.0	12.0	12.0	12.0	12-1	01.04.07 phosphate transport
P060397	strong similarity to novobiosyl 2-dehydrogenase ynfS - Bacillus subtilis	2e-54	AP003185	gene: "CPE0903"; product: "probable dehydrogenase"; Clostrid	13.3	12.0	36.3	93.9	11-13	01.01.10.04 degradation of amino acids of the pyruvate family
P060398	strong similarity to polyamine transport protein Pto1 - Saccharomyces cerevisiae	3e-29	BX49807	gene: "mtdR"; product: "mtdR"; family multistep resistance pro	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060399	strong similarity to transcription factor AtrA - Aspergillus fumigatus	1e-09	SCAR091	product: "transcription factor AtrA"; Aspergillus fumigatus	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060400	strong similarity to polyamine transport protein AA47250.1 - Drosophila melanogaster	3e-24	AE003464	product: "CG3663.PA"; Drosophila melanogaster chromosome	85.2	199.3	180.1	573.8	48-9	01.01.01.01.02 biosynthesis of the glutamate group (proline, hydroxyproline, arginine, glutamine, glutamate)
P060401	strong similarity to sulfur disorganizer like protein AnG040670 - Aspergillus niger	3e-57	AY135619	Gordon's sp. CY252 putative disorganizers, dithienothioam	55.9	40.9	58.6	348.5	0-3	01.05.04 regulation of C-compound and carbohydrate utilization
P060402	strong similarity to alcohol dehydrogenase adhP - Pseudomonas aeruginosa	1e-108	T3387	alcohol dehydrogenase adhP - fission yeast (Schizosaccharomyces	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060403	strong similarity to bile acid transporter Ynf1 - Saccharomyces cerevisiae	1e-06	S64800	probable membrane protein YLL046 - yeast (Saccharomyces c	16.9	30.8	72.8	96.9	10-19	01.01.10.04 degradation of amino acids of the pyruvate family
P060404	strong similarity to hypothetical protein 1144, scaffold, 2 aa, 20aa - Fusarium graminearum	1e-136	SC093810	gene: "SC003711"; "SCF.41.300"; product: "putative secreted pr	12.0	12.0	12.0	12.0	12-1	01.02.01 nitrogen and sulfur utilization
P060405	strong similarity to hypothetical protein PA1077 - Pseudomonas aeruginosa	1e-120	AF493222	product: "hypothetical protein"; Aspergillus fumigatus	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060406	strong similarity to membrane protein Pto1 - Saccharomyces cerevisiae	4e-89	NC8191C9	gene: "Pto1.030"; product: "related to fluoronase resistance pro	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060407	strong similarity to acyl CoA dehydrogenase adhA - Escherichia coli	1e-74	NC2424.2	gene: "ZEA4.020"; product: "conserved hypothetical protein"; Ne	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060408	strong similarity to succinyl-CoA dehydrogenase adhA - Saccharomyces cerevisiae	2e-72	AF003728	product: "GSD55.030"; product: "conserved hypothetical protein";	38.8	68.8	104.8	104.8	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060409	strong similarity to ferulic acid esterase fAseA - Aspergillus niger	1e-130	AF274232	product: "alkaline lipase"; Penicillium cyclopium alkaline lipase r	55.7	144.7	137.2	591.2	86-61	01.04.07 phosphate transport
P060410	strong similarity to hypothetical protein AnG026940 - Aspergillus niger	4e-03	AY048436	product: "putative long-chain acyl-CoA synthetase"; Arabidops	74.5	90.9	256.6	439.0	16-36	01.05.04 regulation of C-compound and carbohydrate utilization
P060411	strong similarity to methylCoA dehydrogenase adhA - Escherichia coli	1e-75	AE018686	product: "nucleotide-binding compound acyl-CoA synthetase. DazA family; I	40.9	78.1	278.8	81.4	76-59	01.05.04 regulation of C-compound and carbohydrate utilization
P060412	strong similarity to hypothetical sterigmatocystin biosynthesis monooxygenase stcV - Aspergillus nidulans	1e-101	BX482595	Neurospora crassa DNA linkage group I BAC clone B24N4.	12.0	12.0	12.0	12.0	12-1	01.01.09 amino acid metabolism activities
P060413	strong similarity to hypothetical short chain dehydrogenase HGT3.130 - Schizosaccharomyces pombe	1e-121	NC55E.16	gene: "frx1B308.100"; product: "related to multistep resistant pro	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060414	strong similarity to integral membrane protein Pto1 - Saccharomyces cerevisiae	1e-130	AF493222	product: "hypothetical protein"; Aspergillus fumigatus	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060415	strong similarity to hypothetical oxoacidase PA0147 - Pseudomonas aeruginosa	2e-48	AP003010	gene: "m15982"; Mesorhizobium loti DNA, complete genome, s	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060416	strong similarity to hypothetical protein PA1213 - Pseudomonas aeruginosa	1e-138	NC51659	gene: "m15982"; product: "conserved hypothetical protein"; Ne	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060417	strong similarity to alantoinase permease AtrA - Aspergillus fumigatus	1e-24	T41694	product: "membrane transport protein - fission yeast (Schizosacchar	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060418	strong similarity to alkane-inducible cytochrome P450 gene ALK1 - Yarrowia lipolytica	1e-133	AB101038	gene: "ALK1"; product: "ALK1"; Yarrowia lipolytica gene for ALK	96.2	133.8	219.7	961.0	10-3	01.05.04 regulation of C-compound and carbohydrate utilization
P060419	strong similarity to carbonyl reductase YnfP - Candida albicans	4e-48	CBY1	product: "CARBOXYLREDUCTASE YnfP [imported] - Candida albicans	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060420	strong similarity to alcohol dehydrogenase adhP - Pseudomonas aeruginosa	1e-108	SC093810	gene: "SC003711"; "SCF.41.300"; product: "putative secreted pr	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060421	strong similarity to L-lysine permease YnfP - Escherichia coli	1e-73	BX1394	product: "lysine permease YnfP"; Escherichia coli	44.9	73.8	105.6	210.0	12-43	01.04.01 phosphate utilization
P060422	weak similarity to enzyme reductase of the flavonoid biosynthesis lovC - Aspergillus terreus	7e-07	AF141925	gene: "lovC"; product: "enzyme reductase"; Aspergillus terreus lov	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060423	strong similarity to hypothetical protein PA1077 - Pseudomonas aeruginosa	1e-120	AF493222	product: "hypothetical protein"; Aspergillus fumigatus	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060424	strong similarity to glycosylphosphatidylinositol-anchored beta1-3glucanase/transferase gbt1 - Aspergillus niger	1e-136	AY105452	unlabeled ORF. Sequence 34 from Patent WO/950303. DP86FC	17.6	114.1	233.1	548.6	12-17	01.02.01.09.09 other catabolism of nitrogenous compounds
P060425	strong similarity to arylsulfatase arfS1 - Neurospora crassa	1e-06	NC077104	DNA centromeric region sequence from BAC DAP1503. DP86FC	155.5	443.0	639.2	1949.6	11-15	01.02.01.09.09 other catabolism of nitrogenous compounds
P060426	strong similarity to glucidase 1A - Aspergillus fumigatus	1e-102	AY1344	product: "glucidase 1A"; Aspergillus fumigatus	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060427	strong similarity to polyamine transport protein Pto1 - Saccharomyces cerevisiae	1e-130	BX49807	gene: "mtdR"; product: "mtdR"; family multistep resistance pro	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060428	strong similarity to hypothetical protein AnG026940 - Aspergillus niger	4e-03	AY048436	product: "putative long-chain acyl-CoA synthetase"; Arabidops	74.5	90.9	256.6	439.0	16-36	01.05.04 regulation of C-compound and carbohydrate utilization
P060429	strong similarity to lactonase-specific esterase estF1 - Pseudomonas aeruginosa	1e-112	SC093911	gene: "SC003711"; "SC003711"; product: "putative monooxygen	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060430	strong similarity to glucose transporter Suc2 - Neurospora crassa	2e-63	NC8188.9	gene: "B888.100"; product: "related to sugar transport protein S	116.3	124.4	62.1	503.5	5-76	01.05.04 regulation of C-compound and carbohydrate utilization
P060431	strong similarity to sugar transporter Suc2 - Pichia stipitis	4e-42	BX482881	gene: "B134.000"; product: "probable maltose permease"; Ne	12.0	12.0	12.0	12.0	12-1	01.05.01 C-compound and carbohydrate utilization
P060432	strong similarity to alpha-glucosidase adhA - Pseudomonas aeruginosa	2e-72	T3387	alcohol dehydrogenase adhP - fission yeast (Schizosaccharomyces	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060433	strong similarity to alpha-amylase precursor amyA - Bacillus amyloliquefaciens	1e-177	AB078784	product: "hypothetical protein"; Aspergillus oryzae gene for hyp	101.5	288.8	265.7	823.1	15-18	01.05.02 C-compound and carbohydrate metabolism
P060434	strong similarity to hypothetical sterigmatocystin biosynthesis monooxygenase stcV - Aspergillus nidulans	1e-116	BX482629	gene: "BUD3.0100"; product: "related to sterol monooxygenase	12.0	12.0	12.0	12.0	12-1	01.01.10 amino acid degradation (catabolism)
P060435	strong similarity to Pto1 - Saccharomyces cerevisiae	1e-130	AF493222	product: "hypothetical protein"; Aspergillus fumigatus	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060436	strong similarity to oxoacidase Pto1 - Saccharomyces cerevisiae	4e-80	NC8188.9	gene: "B888.100"; product: "related to noncarboxylesterase trans	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060437	strong similarity to oxoacidase Pto1 - Saccharomyces cerevisiae	1e-118	NC8188.9	gene: "B888.100"; product: "related to noncarboxylesterase trans	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060438	strong similarity to beta-transtransferase hvt-1 - Pseudomonas aeruginosa	1e-135	AF493222	product: "hypothetical protein"; Aspergillus fumigatus	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060439	strong similarity to lactonase-specific esterase estF1 - Pseudomonas aeruginosa	1e-112	SC093911	gene: "SC003711"; "SC003711"; product: "putative monooxygen	12.0	12.0	12.0	12.0	12-1	01.05.04 regulation of C-compound and carbohydrate utilization
P060440	strong similarity to beta-galactosidase lacA - Aspergillus niger [putative frameshift]	1e-13								

Pc2dg10850	similarly to hypothetical protein An11g00150 - Aspergillus niger	8e-12	T01145	probable acetone-cyanohydrin lyase [imported] - Arabidopsis th	12.6	12.0	25.0	77.3	0	
Pc2dg13360	weak similarity to cytidine deaminase - Homo sapiens	6e-36	S41571	5-hydroxy-3-oxoheptanoate 5,5-lyase (EC 3.5.4.2) - Aspergillus terreus	12.6	12.0	25.0	96.2	0	
Pc2dg13320	weak similarity to hypothetical protein SPB2248.02 - Schizosaccharomyces pombe	1e-09	T40983	2-oxoglutarate 5-lyase (EC 4.1.3.12) - fission yeast (Schizosacchar	22.0	21.2	92.1	140.4	0	
Pc2dg14890	weak similarity to 2-haloacetal halohydrin lyase IVa - Pseudomonas cepacia	1e-05	AP005962	gen: "dmbB"; product: "2-haloacetalic acid dehalogenase"; Bra	12.0	13.3	47.4	243.4	7e-99	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc2dg14920	similarly to hypothetical oxidoreductase P45309 - Pseudomonas aeruginosa	1e-36	NC5F3_10	gen: "SF3.100"; product: "conserved hypothetical protein"; Ne	12.0	12.0	25.0	96.2	0	
Pc2dg14970	weak similarity to hypothetical membrane protein YJ0237v - Saccharomyces cerevisiae	0	0	0	60.8	214.3	143.8	658.1	0	
Pc2dg15400	strong similarity to hypothetical protein hpj0295 - Helicobacter pylori	6e-09	AE018871	product: "polysaccharide deacetylase family protein"; Pseudom	17.1	17.3	47.9	103.9	1e-134	01.01.10.04 degradation of amino acids of the pyruvate family
Pc2dg15440	strong similarity to peroxisomal membrane protein Pmp27 - Saccharomyces cerevisiae	2e-57	AX052571	Unnamed ORF; Sequence 1 from Patent WO0071579.	17.8	36.8	46.2	123.8	4e-39	01.04.07 phosphate transport
Pc2dg15480	strong similarity to hypothetical protein An04g04470 - Aspergillus niger	0	0	0	23.6	59.1	106.1	243.1	0	
Pc2dg15640	strong similarity to hypothetical protein - Bradyrhizobium japonicum	1e-138	AP005959	gen: "bIs8791"; Bradyrhizobium japonicum USDA 110 DNA, cor	12.0	12.0	37.1	105.0	0	
Pc2dg15660	strong similarity to steroid carrier protein-X steroid carrier protein-2 SC2P2 - Homo sapiens	1e-171	NC80811	gen: "B9B81.130"; product: "probable steroid carrier protein"; N	12.0	12.0	23.6	107.1	1e-108	01.07 metabolism of vitamins, cofactors, and prosthetic groups
Pc2dg15670	strong similarity to NAD-GSH-dependent formaldehyde dehydrogenase (H4 - Paracoccus denitrificans	3e-62	B08955	conserved hypothetical protein [imported] - Schizosaccharom	12.0	12.0	49.0	119.1	5e-91	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc2dg160870	strong similarity to vacuolar aminopeptidase Y Ap63 - Saccharomyces cerevisiae	1e-125	BX08808	gen: "G2184.310"; product: "related to aminopeptidase Y pre	27.6	285.5	537.8	1983.1	1e-107	06.01 protein folding and stabilization
Pc2dg17120	similarly to hypothetical neutral amino acid permease - Neurospora crassa	4e-13	SA7892	neutral amino acid permease - Neurospora crassa	12.0	12.4	27.1	104.0	0	
Pc2dg172640	strong similarity to hypothetical protein conig1_1_part_1ts_4603 - Aspergillus fumigatus	0	0	0	47.5	45.3	78.6	213.4	2e-57	99 UNCLASSIFIED PROTEINS
Pc2dg194130	similarly to hypothetical protein SC1C3.21 - Streptomyces coelicolor	5e-11	T34713	hypothetical protein SC1C3.21 - Streptomyces coelic	215.5	369.3	744.2	1317.5	6e-63	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc2dg194310	strong similarity to hypothetical protein SPAC1039.06 - Schizosaccharomyces pombe	1e-74	T50056	hypothetical protein SPAC1039.06 [imported] - fission yeast (Sc	12.0	12.0	38.1	90.1	5e-92	01.01.15 biosynthesis of the pyruvate family (alanine, isoleucine, leucine, valine) and D-alanine
Pc2dg195760	similarly to integral membrane protein PTH11 - Magnaporthe oryzae	1e-120	BX849656	product: "integral membrane protein, putative"; Aspergillus fum	12.0	14.5	12.0	82.3	3e-77	01 METABOLISM
Pc2dg195770	strong similarity to cytochrome P450 eln2 - Coprinus chereus	0	BX849605	product: "cytochrome p450, putative"; Aspergillus fumigatus B4	12.0	25.7	27.9	89.1	1e-128	11.07.01 detoxification involving cytochrome P450
Pc2dg195900	strong similarity to hypothetical short chain dehydrogenase SPAC251.03 - Schizosaccharomyces pombe	1e-110	BX897675	gen: "B5E7.120"; product: "conserved hypothetical protein"; N	144.9	291.9	404.8	779.3	6e-83	01.01.07.06 biosynthesis of lysine
Pc2dg195930	strong similarity to benzate 4-monooxygenase CytH17 - Aspergillus niger	3e-40	AF169116	gen: "CypH"; product: "cytochrome P450 monooxygenase"; Asp	12.0	12.5	32.1	91.9	2e-97	01.05.01 C-compound and carbohydrate utilization
Pc2dg196600	strong similarity to tauroine dihydroase, 2-oxoglutarate-dependent tauD - Escherichia coli	3e-33	AE016756	gen: "tauD"; product: "Alpha-keoglutarate-dependent tauroine	12.0	12.0	12.0	53.1	0	01.01 amino acid metabolism
Pc2dg198020	strong similarity to alcohol dehydrogenase alcB - Aspergillus nidulans	1e-159	SE2746	alcohol dehydrogenase (EC 1.1.1.1) B - Emmericella nidulans	77.1	41.3	154.6	381.8	4e-91	01.07.01 biosynthesis of vitamins, cofactors, and prosthetic groups
Pc2dg198300	strong similarity to gamma-glutamyl transpeptidase GOT1 - Homo sapiens	1e-150	T49441	gamma-glutamyltransferase related protein [imported] - Neurosp	12.0	12.5	89.1	1e-160	01.05.01 C-compound and carbohydrate utilization	
Pc2dg199410	strong similarity to monooamine oxidase maon - Aspergillus niger	0	SE5273	amine oxidase (flavin-containing) (EC 1.4.3.4) N - Aspergillus ni	12.0	26.9	12.0	167.3	1e-107	98 CLASSIFICATION NOT YET CLEAR-CUT
Pc2dg199430	weak similarity to peroxisomal transporter Ant1 - Saccharomyces cerevisiae	1e-42	SE8019	hypothetical protein YPR128c - yeast (Saccharomyces cerevisia	20.4	30.2	48.3	125.3	4e-48	01.05.07 C-compound, carbohydrate transport
Pc2dg199940	strong similarity to hypothetical protein conig5_part_1ts_34203 - Aspergillus fumigatus	4e-97	BX088089	gen: "Z9E9.120"; product: "related to 3-hydroxyisovalerate de	19.7	29.9	54.6	232.3	0	
Pc2dg199950	strong similarity to prokase - Aureobacterium esterarcum	1e-128	NC642_2	gen: "64c2.200"; product: "conserved hypothetical protein"; N	13.7	28.1	75.7	168.1	1e-117	08 CELLULAR TRANSPORT AND TRANSPORT MECHANISMS
Pc2dg121660	strong similarity to flavocytochrome b2 L-lactate dehydrogenase CVB2 - Pichia anomala	1e-3e8	BX443323	gen: "cyb2"; product: "cytochrome b2"; Kluyveromyces lactis cy	12.0	12.0	12.0	52.0	2e-62	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc2dg121980	strong similarity to hypothetical protein conig_1_135_scaffold_11_fta_530w - Aspergillus nidulans	1e-110	BX842620	gen: "B11E5.430"; product: "related to dihydrodipicolinate s	12.0	12.0	14.7	129.4	0	
Pc2dg122200	similarly to hypothetical protein M6r1 - Arabidopsis thaliana	1e-110	AE016757	product: "hypothetical protein"; Escherichia coli C	12.0	12.0	14.7	129.4	0	
Pc2dg133940	strong similarity to hypothetical protein An03g01270 - Aspergillus niger	0	0	0	64.1	74.2	151.9	306.8	1e-118	99 UNCLASSIFIED PROTEINS
Pc2dg14250	strong similarity to hypothetical protein SPAC3H10.01 - Schizosaccharomyces pombe	7e-67	SS5479	hypothetical protein SPAC3H10.01 - fission yeast (Schizosaccha	22.0	85.4	240.3	1064.1	2e-93	01.04 phosphate metabolism
Pc2dg14780	strong similarity to hypothetical protein Anr1g01890 - Aspergillus niger	0	0	0	12.0	12.0	12.0	103.1	0	
Pc2dg15140	strong similarity to 2,4-dihydroxyhept-2-ene-1,7-dioic acid aldolase hpcH - Escherichia coli	4e-44	BX849605	product: "2,4-dihydroxyhept-2-ene-1,7-dioic acid aldolase"; Asp	21.0	22.0	40.1	98.7	7e-65	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc2dg154110	strong similarity to hypothetical protein An07g04880 - Aspergillus niger [putative sequencing error]	4e-06	AS3731	transition initiation factor eIF-2 alpha chain kinase (EC 2.7.1.-)	14.5	16.6	68.2	86.9	0	
Pc2dg15840	strong similarity to NAD2-dependent 15-hydroxyprostaglandin dehydrogenase PGCH - Homo sapiens	8e-18	SC5G2121	gen: "cye-56565"; product: "similar to hydroxyprostaglandin de	38.1	158.0	572.3	367.3	3e-73	11.05.01 resistance proteins
Pc2dg17020	strong similarity to hypothetical protein An04g08050 - Aspergillus niger	0	0	0	12.0	12.0	18.1	64.0	0	
Pc2dg17030	strong similarity to hypothetical protein An04g08060 - Aspergillus niger	0	0	0	12.0	16.6	25.0	72.9	4e-63	01.03 nucleotide metabolism
Pc2dg17620	weak similarity to hypothetical protein TM11177 - Thermoglossa maritima	3e-08	AE017006	product: "hydrolase (HAD superfamily)"; Bacillus cereus ATCC	34.0	28.0	105.0	228.1	0	
Pc2dg18260	weak similarity to hypothetical protein YLL056c - Saccharomyces cerevisiae	1e-06	S50964	hypothetical protein YLL056c - yeast (Saccharomyces cerevisia	37.5	53.5	128.2	190.4	1e-166	99 UNCLASSIFIED PROTEINS
Pc2dg18570	strong similarity to isochromane pyruvate-lyase - Pseudomonas syringae	3e-09	AE016865	gen: "pobB"; product: "isochromane pyruvate-lyase"; Pseudo	25.5	69.3	104.3	169.9	2e-68	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc2dg18580	strong similarity to p-sulfonal alcohol dehydrogenase 1 - Comamonas testosteroni	1e-10	AL846883	gen: "R523251"; "R520195"; product: "PROBABLE TC	12.0	12.0	12.0	53.1	1e-105	01.01.07 amino acid transport
Pc2dg18680	strong similarity to 1,4-butanediol diacylate esterase BDA1 - Brevibacterium lactens	7e-51	BX572594	product: "putative esterase"; Rhodospirillum rubrum	116.6	310.5	680.8	4e-88	01.06 lipid, fatty-acid and isoprenoid metabolism	
Pc2dg19460	strong similarity to hypothetical sterol transporter YMR034c - Saccharomyces cerevisiae	1e-57	SE3591	probable membrane protein YMR034c - yeast (Saccharomyces	12.0	21.4	24.6	94.8	2e-76	01 METABOLISM
Pc2dg195990	strong similarity to hypothetical protein conig1_1_part_1ts_34203 - Aspergillus fumigatus	1e-110	AE3150	hypothetical protein YK1025 [imported] - Agrobacterium tumefaci	12.0	12.0	12.0	53.1	0	
Pc2dg196160	strong similarity to hypothetical protein 1329_scaffold_5_fta_150g - Fusarium graminearum	3e-26	E70900	probable lipase - Mycobacterium tuberculosis (strain H37RV)	39.5	102.4	26.9	323.9	1e-99	01.01.01.15 biosynthesis of the pyruvate family (alanine, isoleucine, leucine, valine) and D-alanine
Pc2dg20520	strong similarity to hypothetical protein conig1469_fta_1450g - Aspergillus fumigatus	1e-11	AX186125	Sequence 197 from Patent WO0151639.	12.0	21.4	83.7	129.0	0	
Pc2dg20910	strong similarity to hypothetical protein conig1469_fta_1450g - Aspergillus fumigatus	1e-145	AF14242	gen: "A1A1242.02"; product: "putative 3-hydroxyisobutyrate de	12.0	12.0	12.0	53.1	0	
Pc2dg20960	strong similarity to lysine-specific permease lysP - Escherichia coli	0	AF11292	gen: "A1A1242.02"; product: "putative lysine-specific permease	68.8	115.0	17.7	42.8	0	
Pc2dg123120	strong similarity to sequence 1 - unknown organism	5e-72	AE100446	Sequence 1 from Patent WO0211779.	12.0	15.0	24.8	65.1	7e-81	01.01.01.07.06 biosynthesis of lysine
Pc2dg12520	similarly to C-6 sterol isomerase - Arabidopsis thaliana	8e-31	AS6122	ompam-binding protein - guinea pig	12.0	12.0	12.0	53.1	0	
Pc2dg12740	strong similarity to tauroine dihydroase, 2-oxoglutarate-dependent like protein An13g02700 - Aspergillus niger [p	0	0	0	12.0	12.0	12.0	53.1	9e-99	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc2dg12950	strong similarity to high-affinity nicotinic acid permease Tna1 - Saccharomyces cerevisiae	1e-169	NC642_1	gen: "64c2.200"; product: "related to PUTATIVE TARTRATE T	12.0	14.1	17.8	80.9	1e-97	01.20.35.01 biosynthesis of phenylpropanoids
Pc2dg122440	strong similarity to cholesterol 7alpha-monooxygenase CYP7 - Oryzotrygon culicoides	2e-17	AF127050	gen: "CYP7B1"; product: "bacterial 7alpha-hydroxylase"; G	12.0	12.0	12.0	53.1	0	
Pc2dg122770	strong similarity to cytochrome P450 monooxygenase YK1025 - Agrobacterium tumefaciens	1e-119	BFX4501	gen: "P450"; product: "cytochrome P450 monooxygenase"; G	31.5	12.0	51.7	116.1	1e-153	01.01.01 amino acid biosynthesis
Pc2dg20100	strong similarity to allantoinase permease Dafs - Saccharomyces cerevisiae	4e-82	BX842634	gen: "B1E68.030"; product: "related to allantoinase permease"; I	12.0	12.0	12.0	37.2	1e-52	01.04.07 phosphate transport
Pc2dg20120	strong similarity to hypothetical protein p010417.1 - Magnaporthe oryzae	0	0	0	57.2	121.6	36.1	849.1	1e-171	01.02.01 nitrogen and sulfur utilization
Pc2dg20220	strong similarity to hypothetical protein conig1770g - Aspergillus niger	4e-46	AX382626	gen: "B11E5.430"; product: "related to dihydrodipicolinate s	12.0	12.0	12.0	53.1	0	
Pc2dg20290	strong similarity to ATP-hydrolyzing 5-oxoprolinase - Rattus norvegicus	0	T37524	probable oxoprolinase - fission yeast (Schizosaccharomyces po	223.3	254.2	1089.9	1649.8	1e-120	01 METABOLISM
Pc2dg20570	strong similarity to triacylglycerol lipase Ipl1 - Geotrichum candidum	2e-42	SA1091	triacylglycerol lipase (EC 3.1.1.3) [precursor - yeast (Geotrichu	85.2	249.2	257.5	704.0	0	
Pc2dg20840	strong similarity to 3-isopropylmalate dehydrogenase icu28 - Aspergillus niger	1e-153	T27210	3-isopropylmalate dehydrogenase (EC 1.1.1.85) B - Aspergillus	15.6	46.7	46.0	327.0	0	
Pc2dg21110	strong similarity to urea transport protein Dur3 - Saccharomyces cerevisiae	1e-138	T39959	probable urea active transporter - fission yeast (Schizosacchar	12.0	12.0	12.0	60.0	0	
Pc2dg21150	strong similarity to nitroimidazole monooxygenase component A - Mesorhizobium loti	0	AP003011	gen: "nirT299"; product: "nitroimidazole monooxygenase comp	37.8	117.3	157.8	802.8	5e-96	01.05.01 C-compound and carbohydrate utilization
Pc2dg21360	strong similarity to tartrate transport protein tnaB - Agrobacterium vitis	1e-174	NC5F3_25	gen: "SF3.250"; product: "conserved hypothetical protein"; Ne	24.1	34.2	23.4	72.5	4e-53	01.05.07 C-compound, carbohydrate transport
Pc2dg21670	strong similarity to hypothetical oxidoreductase SPAC877.14c - Schizosaccharomyces pombe	1e-103	T50285	probable oxidoreductase [imported] - fission yeast (Schizosacch	31.3	80.7	173.8	359.3	0	
Pc2dg27690	weak similarity to AK-coin regulating protein AKR1 - Alternaria alternata	0	0	0	31.3	43.3	70.9	188.8	0	01.01.01 amino acid biosynthesis
Pc2dg28520	weak similarity to heparanase - Homo sapiens	1e-108	SE5273	amine oxidase (flavin-containing) (EC 1.4.3.4) N - Aspergillus ni	22.0	85.4	240.3	1064.1	1e-95	01.01.01.11.09 biosynthesis of phenylalanine
Pc2dg28560	strong similarity to fluconazole resistance transporter Flr1 - Saccharomyces cerevisiae	1e-124	BX088112	gen: "G17A4.250"; product: "related to multidrug resistant por	12.0	23.6	19.9	54.3	2e-53	01.05.07 C-compound, carbohydrate transport
Pc2dg289210	strong similarity to alpha-glucosidase - Bacillus thermoamyloquelacens	0	SC039310	gen: "SC01055"; "SC0221.03"; "SC020A.35c"; product: "puta	12.0	13.8	51.3	98.0	4e-59	01.05.07 C-compound, carbohydrate transport
Pc2dg29220	strong similarity to hypothetical protein conig1813.1 - Neurospora crassa [putative sequencing error]	1e-12	AE011062	gen: "MA353.1"; product: "methylase"; Methanococcus thermo	12.0	12.0	61.5	123.1	3e-70	01.20.39 other secondary metabolic activities
Pc2dg29680	similarly to hypothetical protein PA420A - Pseudomonas aeruginosa	7e-17	AE016869	product: "conserved hypothetical protein"; Pseudomonas syring	226.1	224.8	575.4	1150.1	0	
Pc2dg121250	strong similarity to oligopeptide transporter like protein An15g07460 - Aspergillus niger	5e-77	AY054590	gen: "Z97341.3"; product: "Unknown protein"; Arabidopsis thal	12.0	12.0	25.2	72.7	0	
Pc2dg13360	strong similarity to dihydroxy-acid dehydratase like protein An07g08870 - Aspergillus niger	0	BX840427	gen: "tdD"; product: "putative dihydroxy-acid dehydratase"; B	12.0	25.4	27.2	554.2	0	
Pc2dg13440	strong similarity to hypothetical protein conig1483_1_fta_360w - Aspergillus fumigatus	5e-26	AE14794	polyketide biosynthesis associated protein An1775 [imported];	27.1	79.0	72.1	169.8	0	
Pc2dg13600	strong similarity to hypothetical protein SPAC322.05c - Schizosaccharomyces pombe	9e-41	T50270	hypothetical protein SPAC322.05c [imported] - fission yeast (Sc	12.0	12.0	49.0	375.3	1e-121	99 UNCLASSIFIED PROTEINS
Pc2dg14520	strong similarity to allantoinase permease Dafs - Saccharomyces cerevisiae	4e-99	T41604	probable membrane transport protein - fission yeast (Schizosac	12.0	12.0	14.6	129.1	1e-157	01.01.01.15 biosynthesis of the pyruvate family (alanine, isoleucine, leucine, valine) and D-alanine
Pc2dg14710	strong similarity to bifunctional xylose-4-epimerase xylB - Thermotoga aerotolerans	1e-148	AF135015	gen: "xylB"; product: "xylose-4-epimerase"; Thermotoga	12.0	12.0	12.3	58.0	0	
Pc2dg15100	strong similarity to dicarboxylic amino acid permease Dip5 - Saccharomyces cerevisiae	1e-109	UF430825	gen: "AAT1"; product: "amino acid transporter"; Uromyces fab	186.2	281.1	233.6	421.1	1e-105	01.01.07 amino acid transport
Pc2dg15170	strong similarity to propionyl-CoA carboxylase beta chain precursor like protein An15g02830 - Aspergillus	5e-48	BX872604	product: "possible methylmalonyl-CoA decarboxylase alpha cha	12.0	12.0	14.6	129.1	1e-148	01 METABOLISM
Pc2dg15180	strong similarity to pyruvate carboxylase like protein An15g02820 - Aspergillus niger	4e-93	AB3978	pyruvate carboxylase pycA [imported]; Bacillus halodurans (str	12.0	44.1	31.8	297.1	6e-84	01 METABOLISM
Pc2dg16410	strong similarity to long-chain-fatty-acid acyl-CoA ligase like protein An07g05210 - Aspergillus niger	1e-163	BX842641	gen: "B2221.020"; product: "related to long-chain fatty-acid-C	22.3	84.2	54.8	291.9	1e-126	01 METABOLISM
Pc2dg17460	strong similarity to cytochrome P450 piazin demethylase PDAT9 - Nectria haematococca	3e-50	SA5583	piazin demethylase (EC 1						

Supplementary Table 12. K-mean cluster 4
@=Values given are the average of three independent experiments

orf code	Description of putative <i>P. chrysogenum</i> ORF	Best blast homolog to putative <i>P. chrysogenum</i> ORF	Average transcript levels @				FunCat (auto)	category	
			WIS - PAA	WIS + PAA	DS - PAA	DS + PAA			
p-value	Gene	description							
Pc12g00830	strong similarity to sorbitol utilization protein sou2 - <i>Candida albicans</i>	1e-125 AF307300, gene: "MDH", product: "NADP-dependent mannitol dehydratase"	65.6	91.0	1200.5	757.5	0		
Pc12g01900	strong similarity to nuclear migration factor nufF - <i>Aspergillus nidulans</i>	1e-118 NCBI7423, gene: "B7H23.150", product: "related to NUCLEAR MRGAT"	126.7	126.7	69.9	2e-40	01.01.04 regulation of amino acid metabolism	0	
Pc12g01930	similarity to ribonuclease II like protein An03g0790 - <i>Aspergillus niger</i>	0 0	73.9	73.9	42.3	39e-1	06.04 protein targeting, sorting and translocation	0	
Pc12g05550	weak similarity to fructosamine-3-kinase FNK - <i>Homo sapiens</i>	2e-14 NCBI354, gene: "B3E4.080", product: "conserved hypothetical protein"	16.4	22.1	116.3	16.4	0		
Pc12g07500	strong similarity to alpha-1,3-glucanase muA - <i>Penicillium purpogenum</i>	1e-173 PFU03839g, gene: "muA", product: "alpha-1,3-glucanase", <i>Penicillium</i> fr	17.0	16.3	96.2	130.4	1e-80	01.05.07 C-compound, carbohydrate transport	0
Pc12g07820	similarity to chitinase canA - <i>Aspergillus oryzae</i>	1e-87 AB159785, gene: "canC", product: "chitinase", <i>Aspergillus oryzae</i> c	34.4	26.4	1190.9	1569.5	0		
Pc12g07910	similarity to potassium transport protein Tk2 - <i>Saccharomyces cerevisiae</i>	1e-112 SC020402, gene: "tk1", product: "high-affinity potassium uptake transp	12.0	25.2	87.9	2e-99	01.20 secondary metabolism	0	
Pc12g08710	strong similarity to sugar transport protein S81 - <i>Saccharomyces cerevisiae</i>	1e-44 SG6951, gene: "S81", product: "alpha-1,3-glucanase", <i>Saccharomyc</i> cer	31.3	12.0	134.0	3e-86	01.01.01.15 biosynthesis of the pyruvate family (alanine, isoleucine, leucine, valine) and D-alanine	0	
Pc12g14380	strong similarity to ribonuclease T1 precursor muA - <i>Aspergillus niger</i>	1e-45 A23620, gene: "ribonuclease T1 (EC 3.1.27.3) - <i>Penicillium chrysogenum</i>	12.0	12.0	234.4	72.5	1e-28	01.05 C-compound and carbohydrate metabolism	0
Pc12g14640	strong similarity to hypothetical protein An06g00160 - <i>Aspergillus niger</i>	0 0	98.1	45.7	1520.7	3156.7	0		
Pc13g06200	similarity to pyoverdine biosynthesis protein PvcA - <i>Pseudomonas aeruginosa</i>	2e-23 BX571868, <i>Photobacterium luminescens</i> subsp. laumondii TTO1 compl	22.7	12.2	345.8	35.1	0		
Pc13g07080	similarity to hypothetical protein An15g00820 - <i>Aspergillus niger</i>	0 0	68.9	15.1	678.1	428.5	1e-86	01.05.01 C-compound and carbohydrate utilization	0
Pc13g07880	similarity to 3-hydroxyisobutyrate dehydrogenase mmsB - <i>Pseudomonas aeruginosa</i>	2e-17 AE010190, gene: "PF0716", product: "3-hydroxyisobutyrate dehydroge	12.3	17.3	77.2	134.7	1e-44	01.01.01.11 biosynthesis of the cysteine-aromatic group	0
Pc13g0880	strong similarity to hypothetical protein PA1213 - <i>Pseudomonas aeruginosa</i>	1e-112 NC01509, gene: "G1508.020", product: "conserved hypothetical protein"	67.5	59.0	325.2	367.1	0		
Pc13g11940	strong similarity to glucan 1,4-alpha-glucosidase glaA - <i>Aspergillus niger</i>	1e-134 BX452599, <i>Neurospora crassa</i> DNA linkage group II B4C clone 5204N	91.2	156.0	3281.4	3631.9	1e-111	01.20.17.03 biosynthesis of amines	0
Pc13g14720	strong similarity to pisin demethylase PDA6-1 - <i>Nectria haematococca</i>	7e-35 S34286, pisin demethylase PDA6-1 - fungus <i>Nectria haematococ</i>	12.0	18.4	112.8	164.1	0		
Pc13g14980	weak similarity to developmental protein C-factor csfA - <i>Myxococcus xanthus</i>	4e-20 BX572098, product: "Short-chain dehydrogenase/reductase (SDR) su	93.0	107.4	558.5	336.0	2e-97	01.05.01 C-compound and carbohydrate utilization	0
Pc16g01010	strong similarity to choline monooxygenase like protein An04g09810 - <i>Aspergillus niger</i>	1e-29 AF006580, <i>Gloeobacchar violaceus</i> PCC 1421 DNA, complete genome,	12.0	12.0	80.4	61.1	0.0	01.06.01 lipid, fatty-acid and isoprenoid biosynthesis	0
Pc16g03030	strong similarity to alpha-amyrase precursor amy - <i>Aspergillus niger</i>	0 AB008370, product: "acid-stable alpha-amyrase", <i>Aspergillus kawachi</i>	48.3	68.4	503.4	910.0	4e-58	01.01.01 amino acid biosynthesis	0
Pc16g03010	strong similarity to hypothetical 6-hydroxy-D-nicotine oxidase related protein 13E11.250 - <i>Neurospora crassa</i>	2e-61 T48777, 6-HYDROXY-D-NICOTINE OXIDASE related protein [mpoi	12.0	12.0	112.4	109.1	0		
Pc16g03020	weak similarity to integral membrane protein PTH11 - <i>Magnaporthe grisea</i>	0 0	12.0	12.0	176.4	169.9	0.0	99 UNCLASSIFIED PROTEINS	0
Pc16g07460	strong similarity to hypothetical neutral amino acid permease - <i>Neurospora crassa</i>	1e-143 BX484218, gene: "B482.030", product: "related to neutral amino acid p	32.3	68.2	359.3	411.1	0		
Pc16g10920	strong similarity to hypothetical protein mg07765.1 - <i>Magnaporthe grisea</i>	0 0	21.9	12.0	92.8	90.2	1e-116	01.01.01 amino acid biosynthesis	0
Pc16g11480	strong similarity to lovastatin diolide synthase lovF - <i>Aspergillus terreus</i>	0.0 CHU88040, gene: "PKS1", product: "polyketide synthase", <i>Cochliobolus</i>	12.0	12.0	688.5	208.1	2e-53	01.05.07 C-compound, carbohydrate transport	0
Pc16g11490	weak similarity to hypothetical protein 11f5, scaffold, 2 fts, 130w - <i>Fusarium graminearum</i>	1e-11 OSJN000009, gene: "OSJN000814.2", <i>Oryza sativa</i> genomic DNA	12.0	12.0	201.5	91.8	1e-113	01.20 secondary metabolism	0
Pc16g11500	strong similarity to hypothetical integral membrane protein - <i>Schizosaccharomyces pombe</i>	1e-134 T39090, probable integral membrane protein - fission yeast <i>Schizo</i>	82.1	79.1	644.7	345.2	0		
Pc16g12080	strong similarity to allergen Asp f4 - <i>Aspergillus fumigatus</i>	5e-55 AFASPPF, gene: "rasp f4", product: "Asp f4", <i>Aspergillus fumigatus</i> r	88.0	54.8	2292.5	902.5	5e-76	01.01.07 amino acid transport	0
Pc16g13290	strong similarity to hypothetical protein CP0503 - <i>Chlamydomonas reinhardtii</i>	1e-176 NC13831, gene: "B1383.110", product: "conserved hypothetical protei	12.0	12.0	60.5	77.3	0		
Pc16g15380	weak similarity to hypothetical protein An14g0200 - <i>Aspergillus niger</i>	0 0	32.3	52.4	198.9	198.9	0		
Pc18g03640	similarity to hypothetical protein dag11 - <i>Agaricus bisporus</i>	5e-16 AB217103, gene: "dag11", product: "hypothetical protein", <i>Agaricus bis</i>	40.3	25.3	130.1	204.2	1e-112	99 UNCLASSIFIED PROTEINS	0
Pc20g03900	strong similarity to hypothetical amino transporter SPCC18.02 - <i>Schizosaccharomyces pombe</i>	1e-39 T41145, probable amino transporter - fission yeast <i>Schizosacchar</i>	28.1	62.3	482.7	274.0	2e-94	01.05 C-compound and carbohydrate metabolism	0
Pc20g06780	strong similarity to triphosphate 3-O-acetyltransferase TRH01 - <i>Fusarium sporotrichoides</i>	1e-44 SS5857, probable membrane protein TRH0363 - yeast <i>Saccharomy</i>	12.0	12.0	201.5	91.8	1e-113	01.20 secondary metabolism	0
Pc20g07050	strong similarity to fructose-1,6-bisphosphate aldolase - <i>Thermus aquaticus</i>	1e-72 BX06880, gene: "25E8.130", product: "related to fructose-bisphospha	82.0	12.0	57.4	65.8	0		
Pc20g10040	strong similarity to choline permease Hm1 - <i>Saccharomyces cerevisiae</i> [putative sequencing error]	1e-78 SI1175, choline transport protein - yeast <i>Saccharomyces cerevisi</i>	12.0	12.0	114.9	83.5	0		
Pc20g10750	strong similarity to hypothetical protein An06g01170 - <i>Aspergillus niger</i> [truncated ORF]	0 0	34.2	17.9	315.1	347.9	0		
Pc20g10870	strong similarity to cytochrome P450 trichothecene oxygenase TRH - <i>Fusarium sporotrichoides</i>	1e-100 MRAP4941, gene: "TRH", product: "cytochrome P450", <i>Myrothecium r</i>	47.3	12.0	125.6	166.7	1e-95	01.06 lipid, fatty-acid and isoprenoid metabolism	0
Pc20g15260	strong similarity to glutaminase A glaA - <i>Aspergillus oryzae</i>	1e-107 AB029553, gene: "glaA", product: "glutaminase A", <i>Emmericella nidulans</i>	14.8	16.7	89.0	114.8	2e-93	01.04 phosphate metabolism	0
Pc20g15400	strong similarity to cytochrome P450 monooxygenase P450L - <i>Gibberella fujikuroi</i>	7e-79 GFP450L, gene: "P450L", product: "cytochrome P450 monooxygenase	12.0	12.0	287.0	44.8	2e-71	01.05.04 regulation of C-compound and carbohydrate utilization	0
Pc20g16200	strong similarity to phosphatase-repressible phosphate permease pho-4 - <i>Neurospora crassa</i>	1e-105 J20116, phosphatase-repressible phosphate permease - <i>Neurospora</i>	12.0	12.0	45.0	45.0	0		
Pc20g16760	similarity to hypothetical conserved protein SC03.075 - <i>Streptomyces coelicolor</i>	5e-21 AB071418, gene: "ACMSD", product: "2-amino-3-carboxymuconate-4-	16.9	21.2	134.1	221.1	0		
Pc21g04880	hypothetical protein	0 0	14.7	12.0	115.4	141.2	0		0
Pc21g08490	weak similarity to hypothetical short chain dehydrogenase SPCC736.13 - <i>Schizosaccharomyces pombe</i>	5e-44 AE011685, gene: "XAC0583", product: "oxidoreductase", <i>Xanthomonas</i>	12.0	12.0	61.4	59.0	0		0
Pc21g09220	similarity to hypothetical protein 11f5, scaffold, 2 fts, 130w - <i>Fusarium graminearum</i>	1e-11 OSJN000009, gene: "OSJN000814.2", <i>Oryza sativa</i> genomic DNA	12.0	12.0	201.5	91.8	1e-113	01.01.04 regulation of amino acid metabolism	0
Pc21g09240	strong similarity to signal peptidase subunit like protein An01g0560 - <i>Aspergillus niger</i>	1e-20 BC061447, gene: "MGC76332", product: "MGC76332 protein", <i>Xenopus</i>	12.0	12.0	113.4	124.0	5e-82	01.01.07 amino acid transport	0
Pc21g09270	strong similarity to hypothetical protein B11B22.80 - <i>Neurospora crassa</i>	5e-29 T49806, hypothetical protein B11B22.80 [Imported] - <i>Neurospora cr</i>	12.0	12.0	190.6	345.7	5e-69	01.05.01.01.02 polysaccharide degradation	0
Pc21g09830	strong similarity to high affinity glucose transporter HGT1 - <i>Kluyveromyces fragilis</i>	4e-80 NCBI8312, gene: "HGT1.230", product: "related to hexose transporter	43.4	21.0	184.5	141.9	0		0
Pc21g12340	similarity to hypothetical transcription regulator SPAC139.03 - <i>Schizosaccharomyces pombe</i>	1e-10 BX088903, gene: "25E8.130", product: "conserved hypothetical protei	12.0	12.0	61.4	59.0	6e-41	01.01 amino acid metabolism	0
Pc21g12390	strong similarity to hypothetical protein B2123.120 - <i>Neurospora crassa</i>	2e-62 NC065A3, gene: "G65A3.130", product: "probable DFGS protein", <i>Nec</i>	74.6	65.4	556.6	487.8	0.0	01.20.37.03 biosynthesis of peptide antibiotics	0
Pc21g12570	similarity to hypothetical protein SPAC1305.04 - <i>Schizosaccharomyces pombe</i>	5e-15 AS13228, unnamed ORF: Sequence 1 from Patent WO02063018.	51.1	52.7	1113.7	872.7	0		0
Pc21g12590	similarity to 6-hydroxy-D-nicotine oxidase 6-HNO - <i>Anthracobacter oxidans</i>	1e-15 H2ND, 6-HYDROXY-D-NICOTINE OXIDASE (EC 1.5.3.5) 6-HNO	56.7	103.6	2145.5	1607.9	1e-130	01.01.01 amino acid biosynthesis	0
Pc21g12610	weak similarity to 7alpha-cephem-methoxylase subunit cmcJ - <i>Streptomyces lactamdurans</i>	1e-23 GFU41749, gene: "des", product: "GA4 desaturase", <i>Gibberella fujikuroi</i>	15.2	18.5	567.4	366.5	4e-51	01.05.07 C-compound, carbohydrate transport	0
Pc21g12620	strong similarity to asparagine synthase asrB - <i>Bacillus subtilis</i>	2e-83 BX294156, gene: "asrB", product: "asparagine synthetase", <i>Pirella</i> s	55.4	64.3	1901.5	1359.0	3e-78	01.06 lipid, fatty-acid and isoprenoid metabolism	0
Pc21g12630	similarity to peptide synthase pesA - <i>Metazhizium anisopliae</i>	0 AF46045, gene: "text1", product: "nonribosomal peptide synthetase", <i>E</i>	44.4	15.2	1486.7	792.0	1e-112	99 UNCLASSIFIED PROTEINS	0
Pc21g16190	strong similarity to high-affinity nicotinic acid permease Tra1 - <i>Saccharomyces cerevisiae</i>	3e-53 NC64C2, gene: "64C2.200", product: "related to PUTATIVE TARTRA	94.7	161.0	743.3	176.8	5e-57	01.05.04 regulation of C-compound and carbohydrate utilization	0
Pc21g14620	strong similarity to lysophospholipase like protein An16g0180 - <i>Aspergillus niger</i>	1e-77 AX95297, unnamed ORF: Sequence 5 from Patent WO03097825.	66.1	40.5	250.2	697.8	0		0
Pc21g15310	weak similarity to hypothetical intracellular protease amidase related enzyme of the ThiU family CAC2826 - <i>Neurospora crassa</i>	2e-16 AE016866, product: "ThiU/Plp family protein", <i>Pseudomonas syringae</i>	33.0	12.0	362.0	71.1	0		0
Pc21g15460	strong similarity to maculaen dethioylation protein MAK1 - <i>Nectria haematococca</i>	9e-46 S70702, maculaen dethioylation protein 1 - fungus <i>Nectria haemat</i>	345.6	161.9	1820.6	653.9	1e-65	01 METABOLISM	0
Pc21g17820	strong similarity to hypothetical protein sif1024 - <i>Synechocystis sp.</i>	1e-119 NC64C2, 2-gene: "64C2.210", product: "conserved hypothetical protei	92.7	103.8	738.5	1039.6	2e-76	01 METABOLISM	0
Pc21g17980	strong similarity to hypothetical protein conig5_part.1, 2530cp - <i>Aspergillus fumigatus</i>	0 0	77.9	35.7	886.2	144.5	0		0
Pc21g18040	similarity to conserved domain protein - <i>Pseudomonas syringae</i>	2e-24 AE016873, product: "conserved domain protein", <i>Pseudomonas syring</i>	132.4	77.5	606.4	461.7	0		0
Pc21g18920	similarity to cyanate lyase cysS - <i>Escherichia coli</i>	6e-56 NCBIUN1, gene: "B10N12.080", product: "probable cyanate lyase", <i>N</i>	31.8	16.1	159.5	171.4	0		0
Pc21g1280	strong similarity to methyl sterol oxidase Erg25 - <i>Saccharomyces cerevisiae</i>	1e-89 T38986, probable c-4 methyl sterol oxidase - fission yeast <i>Schizos</i>	44.0	81.5	542.2	971.5	3e-91	01.06 lipid, fatty-acid and isoprenoid metabolism	0
Pc21g1290	strong similarity to hypothetical protein mg02069.1 - <i>Magnaporthe grisea</i>	7e-23 AG3272, probable lactoylglutathione lyase (EC 4.4.1.5) [Imported] - <i>E</i>	119.4	95.7	3398.5	2598.6	0		0
Pc21g1420	strong similarity to hypothetical protein An09g04870 - <i>Aspergillus niger</i>	1e-122 BX484220, gene: "B11E5.350", product: "conserved hypothetical protei	12.0	12.0	87.6	107.9	0		0
Pc21g1440	hypothetical protein	0 0	20.0	12.0	533.7	170.9	0		0
Pc21g1460	hypothetical protein	0 0	121.7	87.8	711.9	614.2	1e-127	01.06.01 lipid, fatty-acid and isoprenoid biosynthesis	0
Pc21g1470	strong similarity to hypothetical protein conig1_1_94, scaffold, 6.1fa, 290wg - <i>Aspergillus nidulans</i>	0 0	112.6	82.1	700.3	694.7	0		0
Pc21g12230	strong similarity to hypothetical protein 11f5, scaffold, 2 fts, 130w - <i>Fusarium graminearum</i>	1e-11 OSJN000009, gene: "OSJN000814.2", <i>Oryza sativa</i> genomic DNA	12.0	12.0	109.6	27.6	0		0
Pc21g22250	strong similarity to T-2 toxin biosynthesis protein TR17 - <i>Fusarium sporotrichoides</i>	9e-22 AF336365, gene: "TR17", product: "TR17", <i>Gibberella zeae</i> 88-1 trichoth	27.9	13.3	168.2	404.8	1e-59	01.01.01.11 biosynthesis of the cysteine-aromatic group	0
Pc21g22260	similarity to cholesterol 7alpha-monooxygenase CYP7 - <i>Oryctolagus cuniculus</i>	6e-14 RRC10U, gene: "CYP7A1", product: "cholesterol 7a-hydroxylase", <i>R</i>	34.9	13.7	106.3	399.9	2e-90	01.01.01 amino acid biosynthesis	0
Pc21g22710	strong similarity to hypothetical neutral amino acid permease - <i>Neurospora crassa</i>	7e-49 SA7892, neutral amino acid permease - <i>Neurospora crassa</i>	12.0	12.0	74.4	148.9	1e-101	14.04.03.05 sporulation and germination	0
Pc21g23810	strong similarity to hypothetical transcriptional regulator CAF32162.1 - <i>Aspergillus fumigatus</i>	8e-31 BX484667, product: "transcriptional regulator, putative", <i>Aspergillus fu</i>	12.7	13.3	467.3	645.1	0.0	08.19 cellular import	0
Pc21g23820	strong similarity to hypothetical protein conig1495_1, 1fa, 1060wg - <i>Aspergillus fumigatus</i>	0 0	32.2	24.4	2288.7	2557.1	0		0
Pc22g01910	strong similarity to hypothetical cell wall protein binB - <i>Aspergillus nidulans</i>	2e-47 AN011296, gene: "binB", product: "putative cell wall protein", <i>Aspergill</i>	40.0	205.9	1384.8	2908.5	3e-50	99 UNCLASSIFIED PROTEINS	0
Pc22g06070	strong similarity to sorbitol utilization protein sou2 - <i>Candida albicans</i>	1e-55 SOUL_CAI5ORBITOL_UTILIZATION PROTEIN SOUL2	12.0	12.0	109.6	27.6	0		0
Pc22g01090	strong similarity to hypothetical protein An1g02070 - <i>Aspergillus niger</i>	0 0	25.1	46.3	631.9	85.5	1e-164	01.05.01 C-compound and carbohydrate utilization	0
Pc22g04940	strong similarity to 2-ketoadonate reductase like protein An01g07030 - <i>Aspergillus niger</i>	1e-96 BX484667, product: "NAD-dependent D-isomer specific 2-hydroxyacid	17.9	30.6	141.2	301.5	1e-110	01.07 metabolism of vitamins, cofactors, and prosthetic groups	0
Pc22g07880	strong similarity to alpha-1,6-glucosidase g4A - <i>Aspergillus parasiticus</i>	0 AB044389, gene: "malT", product: "malinase", <i>Aspergillus oryzae</i> malt	132.9	118.5	719.4	1007.1	1e-151	01.05 C-compound and carbohydrate metabolism	0
Pc22g08110	strong similarity to cytochrome P450 rhb1 - <i>Aspergillus nidulans</i>	8e-32 BX484254, gene: "B11D6.150", product: "related to O-methyltransferase	31.6	12.0	189.5	91.6	2e-71	01 METABOLISM	0
Pc22g12680	weak similarity to hypothetical protein PA5433 - <i>Pseudomonas aeruginosa</i>	2e-05 SC50347, hypothetical protein YR042c - yeast <i>Saccharomyces cere</i>	62.1	67.8	309.2	333.5	7e-61	01.02 nitrogen and sulfur metabolism	0
Pc22g13580	strong similarity to diglycosylase related protein like protein An03g00500 - <i>Aspergillus niger</i>	0.0 THA4382, gene: "P2", product: "P2 protein", <i>Trichoderma harzianum</i>	12.						

Supplementary Table 13. K-mean cluster 5

@ =Values given are the average of three independent experiments

orf code	Description of putative <i>P. chrysogenum</i> ORF	Best blast p-value	homolog to putative <i>P. chrysogenum</i> ORF Gene description	Average transcript levels @				FunCat (auto)	
				WIS - PAA	WIS + PAA	DS - PAA	DS + PAA	pval	category
Pc12g0021	strong similarity to quinate transport protein quidJ - Aspergillus nidulans	2e-50	S08498 quinate transport protein - Emmericella nidulans	400.3	375.9	571.3	1262.1	2e-47	01.04.07 phosphate transport
Pc12g0022	strong similarity to galactoside acetyltransferase lacA - Escherichia coli	1e-33	AL935260, gene: "thgA3"; product: "galactoside O-acetyltransferase"; Lactobacillus	471.6	364.9	544.7	960.3	1e-169	01.01.01 amino acid biosynthesis
Pc12g0025	strong similarity to transmembrane transporter lzt1p - Schizosaccharomyces pombe	1e-71	T40485 transmembrane transporter Lzt1p - fission yeast (Schizosaccharomyces pombe)	17.0	12.0	24.1	32.9	2e-79	01.05 C-compound and carbohydrate metabolism
Pc12g0105	strong similarity to threonine aldolase Gly1 - Schizosaccharomyces pombe	9e-87	AQA15442, gene: "GLY1"; product: "threonine aldolase"; Emmericella nidulans	279.2	401.4	777.3	546.1	0	
Pc12g0114	strong similarity to alcohol dehydrogenase alcB - Aspergillus nidulans	5e-49	AB046446, gene: "alcB"; product: "alcohol dehydrogenase II"; Monascus anka 1	12.0	12.0	12.0	32.0	1e-102	01 METABOLISM
Pc12g0235	weak similarity to phosphotriphosphatase acetyltransferase pat - Streptomyces coelicolor	9e-07	AP005935, gene: "bli0009"; Bradyrhizobium japonicum USDA 11A DNA, compl	21.8	17.1	32.6	58.4	0	
Pc12g0261	strong similarity to 3-phosphoglycerate dehydrogenase like protein An12g01580 - Aspergillus	5e-46	AB3200 phosphoglycerate dehydrogenase serA [imported] - Agrobacterium tu	12.0	12.0	12.0	26.0	8e-75	01.05.07 C-compound, carbohydrate transport
Pc12g0295	weak similarity to 2-hydroxysulfonate reductase IRL - Zea mays	2e-11	NC5F3_3, gene: "5F3.030"; product: "hypothetical protein"; Neurospora crassa	126.3	58.5	152.4	152.6	2e-57	01 METABOLISM
Pc12g0322	strong similarity to high-affinity glucose transporter HGT11 - Kluyveromyces fragilis	1e-84	AF168613, gene: "thgA"; product: "hexose transporter protein"; Aspergillus para	65.9	21.8	30.3	51.4	0.0	01.05.01.01 C-compound, carbohydrate catabolism
Pc12g0351	strong similarity to lipase LipP - Pseudomonas sp.	1e-23	F84000, gene: "lipP"; product: "trichothecene C-15 hydroxylase"; Gibberella	90.2	29.9	159.5	108.4	1e-135	01.01 amino acid metabolism
Pc12g0365	strong similarity to beta-glucosidase bglJ - Candida molischiana	0.0	A46838_1, unnamed ORF; Sequence 5 from Patent WO0553009	36.1	31.9	51.8	117.4	1e-107	01.01 amino acid metabolism
Pc12g0431	strong similarity to phosphoserine transaminase Ser1 - Saccharomyces cerevisiae	1e-121	NCB20D11, gene: "B20D17.010"; product: "related to 3-phosphoserine aminotran	343.4	363.1	806.0	757.1	8e-78	01.05.07 C-compound, carbohydrate transport
Pc12g0545	strong similarity to O-acetylthiomoserine (thiol)-lyase cysD - Aspergillus nidulans	0.0	CYS_D, EM-O-ACETYLTHIOMOSERINE (THIO)-LYASE (EC 4.2.99.10) (O-ACE	697.3	788.1	1322.9	1708.2	0	
Pc12g0544	strong similarity to maltose transport protein Mal31 - Saccharomyces cerevisiae	6e-93	NCSE5_16, gene: "5E6.180"; product: "probable alpha-glucoside transport protei	235.5	236.4	595.9	933.0	1e-120	01.01.10.04 degradation of amino acids of the pyruvate family
Pc12g0831	strong similarity to precursor of tannase - Aspergillus oryzae	0.0	TST50593, gene: "taeC"; product: "ferulic acid esterase"; Talaromyces stipitatus	29.7	37.6	35.6	79.8	2e-80	01.01.01.01 assimilation of ammonia, biosynthesis of the glutamate group
Pc12g0851	strong similarity to long-chain acyl-CoA dehydrogenase like protein An13g03940 - Aspergillus	1e-101	AY033596, product: "probable acyl-CoA dehydrogenase"; Glomus intraradices p	135.7	126.8	178.3	352.8	6e-93	01.01.10 amino acid degradation (catabolism)
Pc12g0884	weak similarity to thyroxine hormone-binding protein mu-crystallin Crn - Homo sapiens	2e-36	ANA3458, gene: "prnX"; product: "PrnX protein"; Aspergillus nidulans prnD, prn	44.7	74.4	79.0	107.1	1e-147	01.01 amino acid metabolism
Pc12g0981	strong similarity to acetamidase amdS - Aspergillus oryzae	0.0	AY048562, gene: "gmdA"; product: "general amidase"; Emmericella nidulans gene	45.7	61.8	166.9	198.5	0	
Pc12g1031	strong similarity to L-2,4-diaminobutyrate decarboxylase dat - Acinetobacter baumannii	1e-104	NC1234A, gene: "1234A.310"; product: "related to L-2,4-diaminobutyrate decar	31.1	30.1	92.4	84.3	5e-49	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc12g1095	strong similarity to secretory lipase LIP2 - Candida albicans	1e-117	AO15710, unnamed ORF; Sequence 1 from Patent WO9954020	93.1	32.4	29.5	107.2	7e-54	01.05 C-compound and carbohydrate metabolism
Pc12g1154	strong similarity to isopenicillin N epimerase cepD - Streptomyces clavuligerus	2e-45	T40624, probable class v pyridoxal phosphate dependent aminotransferase - f	272.6	431.8	805.3	1032.8	1e-108	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc12g1171	strong similarity to beta-glucuronidase GUSB - Canis lupus	2e-82	SC083913, gene: "SCO7506"; "SCBAC17A6.39"; product: "putative hydrolase";	31.2	39.2	112.3	101.4	3e-74	01 METABOLISM
Pc12g1231	strong similarity to multifunctional beta-oxidation protein - Neurospora crassa	1e-104	AY102594, gene: "TRY1"; product: "trichothecene C-15 hydroxylase"; Gibberella	44.3	34.7	57.2	69.4	1e-174	01 METABOLISM
Pc12g1336	strong similarity to aliphatic nitrilase - Rhodococcus rhodocrous	3e-93	NCB23G1, gene: "B23G1.100"; product: "related to aliphatic nitrilase"; Neurosp	18.0	13.8	25.4	40.3	9e-66	01.05 C-compound and carbohydrate metabolism
Pc12g1361	strong similarity to indole-3-acetaldehyde dehydrogenase idt1 - Ustilago maydis	1e-101	T51919, related to aldehyde dehydrogenase (NAD+) [imported] - Neurospora c	213.9	147.6	360.4	464.3	4e-63	01.01 amino acid metabolism
Pc12g1421	strong similarity to enoyl reductase of the lovastatin biosynthesis locV - Aspergillus terreus	2e-42	AF002213, product: "putative dehydrogenase"; Podospora anserina idt-2 locus, t	129.9	179.2	196.2	407.2	0	
Pc12g1461	strong similarity to choline permease Hnm1 - Saccharomyces cerevisiae	3e-57	NI131668, gene: "gabA"; product: "GABA permease"; Emmericella nidulans gabA	46.9	36.6	71.1	114.5	1e-101	01 METABOLISM
Pc13g0251	strong similarity to L-2-haloalkanoic acid dehalogenase hadL - Pseudomonas putida	1e-19	HAD_PSE-2-HALOALKANOIC ACID DEHALOGENASE (EC 3.8.1.2) (L-2-HAL	15.4	15.9	47.8	60.8	3e-39	01.01.01 amino acid biosynthesis
Pc13g0271	strong similarity to glutamate decarboxylase pdc1 - Neurospora crassa	4e-32	BX572603, product: "MacC-like dehydratase Aspergillus glaucus (glutamate)"; Rhod	441.9	416.2	460.4	853.4	1e-65	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc13g0406	strong similarity to hypothetical pyridoxine synthesis protein PDX2 - Cercospora nicotianae	1e-100	AF406113, gene: "pyrB"; product: "pyridoxine"; Aspergillus nidulans pyridoxine	82.7	101.3	164.3	241.5	1e-145	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc13g0411	strong similarity to 7alpha-cephem-methoxylease subunit cmcJ - Streptomyces lactamdurans	1e-49	AP005958, gene: "bli6449"; Bradyrhizobium japonicum USDA 11A DNA, compl	15.4	12.0	12.0	47.8	7e-59	01.01.10 amino acid degradation (catabolism)
Pc13g0441	strong similarity to diacylglycerol acyltransferase DAGAT - Arabidopsis thaliana	9e-63	AF251794, product: "putative diacylglycerol acyltransferase"; Brassica napus c	59.7	46.2	65.9	110.0	1e-141	01.01.99 other amino acid metabolism activities
Pc13g0511	strong similarity to phenol 2-monooxygenase - Trichosporon beigelii	0.0	BX088810, gene: "G3C5.110"; product: "probable phenol 2-monooxygenase"; N	116.8	95.9	195.8	238.7	0.0	01.01.01 amino acid biosynthesis
Pc13g0541	strong similarity to leukotriene-A4 hydrolase - Mus musculus	1e-161	BE10099, leukotriene-A4 hydrolase homolog YNL045w - yeast (Saccharomyces	168.0	167.0	253.6	396.9	0	
Pc13g0551	strong similarity to ATP-dependent 5-oxopropinate - Rattus norvegicus	0.0	T37521, product: "fission yeast (Schizosaccharomyces pombe) 5-oxopropinate	43.7	38.0	101.5	98.1	9e-89	01 METABOLISM
Pc13g0665	strong similarity to C4-dicarboxylate transport protein mae1p - Schizosaccharomyces pombe	1e-70	NCB21D9, gene: "B21D9.140"; product: "related to C4-dicarboxylate transport p	12.0	12.0	13.8	34.0	0.0	01.05 C-compound and carbohydrate metabolism
Pc13g0681	strong similarity to phenazine biosynthesis oxidoreductase phzF - Pseudomonas fluorescens	4e-22	S75237, hypothetical protein - Synchocystis sp. (strain PCC 6803)	213.7	268.4	417.6	610.8	7e-79	01.05.07 C-compound, carbohydrate transport
Pc13g0771	strong similarity to succinate-semialdehyde dehydrogenase NAD(P)+ gabD - Escherichia coli	1e-158	AE016857, gene: "gabD-2"; product: "succinate-semialdehyde dehydrogenase";	30.4	52.8	76.6	120.3	0.0	02.11 electron transport and membrane-associated energy conservation
Pc13g0865	strong similarity to high-affinity glucose transporter HGT1 - Kluyveromyces fragilis	1e-73	A31776, lacase permease - yeast (Kluyveromyces marxianus var. lactis)	85.8	31.3	56.9	116.5	1e-124	01.01.01 amino acid biosynthesis
Pc13g0861	strong similarity to FAD dependent L-sorbose dehydrogenase SDH - Glucanobacter oxydans	7e-53	AP003001, gene: "mh3225"; product: "dehydrogenase; polyethylene glycol dehyd	44.8	55.2	100.0	149.2	1e-150	01.01 amino acid metabolism
Pc13g0881	strong similarity to homoserine dehydrogenase YJR139c - Saccharomyces cerevisiae	1e-109	T40733, probable homoserine dehydrogenase [imported] - Neurospora crassa	325.2	592.6	652.7	871.4	3e-89	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc13g0932	strong similarity to glutamate decarboxylase pdc1 - Neurospora crassa	2e-81	AF149831, gene: "GAD67"; product: "glutamate acid decarboxylase isoform 1"; P	145.5	145.5	249.6	371.5	1e-145	01.03.01 purine nucleotide metabolism
Pc13g0971	strong similarity to fructosyl amino oxygen oxidoreductase - Aspergillus fumigatus	0.0	PJFA001, gene: "facP"; product: "fructosyl amino acid oxidase"; P. janthinellum	48.3	43.4	117.8	193.1	2e-83	01.01.01.15 biosynthesis of the pyruvate family (alanine, isoleucine, leucine, valine) and D-alanine
Pc13g1101	strong similarity to inosine-uridine nucleoside hydrolase IUNH - Crithidia fasciculata	1e-154	BX64905, product: "possible nucleoside hydrolase"; Aspergillus fumigatus BAC	101.0	101.8	201.4	205.8	3e-55	01 METABOLISM
Pc13g1191	strong similarity to 3-hydroxyisobutyrate dehydrogenase mmsB - Pseudomonas aeruginosa	6e-35	BX64906, product: "hypothetical protein"; Aspergillus fumigatus BAC pilot proje	23.7	30.1	43.8	74.1	6e-62	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc13g1201	strong similarity to acetyl-hydrolase chnC - Acinetobacter sp.	2e-87	BX64906, product: "esterase/lipase/cholesterase family protein, putative"; Aspe	111.1	147.1	180.3	440.2	1e-143	01.06.99 other lipid, fatty-acid and isoprenoid metabolism activities
Pc13g1341	strong similarity to hypothetical short-chain dehydrogenase PA1379 - Pseudomonas aeruginosa	2e-36	NCB1181, product: "short-chain oxidoreductase, putative"; Aspergillus fumigatu	90.1	176.9	198.7	223.8	1e-172	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc13g1411	strong similarity to glycosylphosphatidylinositol-anchored beta1-3-glucanase/transferase gel1 - Rattus norvegicus	1e-123	NCB1781, gene: "B1781.110"; product: "related to beta (1-3) glucanase/transfer	64.8	73.7	126.5	184.4	9e-74	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc13g1561	strong similarity to indoleamine 2,3-dioxygenase IDO - Mus musculus	9e-83	S57097, indoleamine-pyrole 2,3-dioxygenase homolog YJR078w - yeast (S	53.3	87.0	127.8	198.8	2e-84	01.05.01.01.01 sugar, glucoside, polyol and carboxylate catabolism
Pc13g1581	strong similarity to serine racemase - Mus musculus	1e-107	NCB11H2, gene: "B11H2.140"; product: "related to threonine dehydratase"; N	56.2	60.5	114.2	136.1	1e-110	01.01.07 amino acid transport
Pc13g1591	strong similarity to endo 1,5-alpha-arabinanase abnA - Aspergillus niger	1e-122	ABNA, ASI ARABINAN ENDO-1,5-ALPHA-L-ARABINOSIDASE A PRECURSOR	131.4	255.9	402.0	547.5	1e-175	01.01 amino acid metabolism
Pc14g0011	strong similarity to lysine permease Lyp1 - Saccharomyces cerevisiae	1e-129	S60914, lysine transport protein - yeast (Saccharomyces cerevisiae)	38.7	14.5	45.6	54.0	4e-58	01.05.07 C-compound, carbohydrate transport
Pc14g0021	strong similarity to aromatic L-amino acid decarboxylase DDC - Rattus norvegicus	1e-103	RN31884, product: "aromatic L-amino acid decarboxylase"; Rattus norvegicus a	76.5	95.8	125.7	224.4	0.0	01 METABOLISM
Pc14g0041	strong similarity to allantoate permease Dals - Saccharomyces cerevisiae	9e-62	BX42680, gene: "90CA.090"; product: "related to allantoate permease"; Neuros	290.2	412.5	470.9	862.1	1e-76	01.04 phosphate metabolism
Pc14g0051	strong similarity to succinyl-CoA:3-ketoad-CoA transferase SCOT - Homo sapiens	1e-131	BT008292, product: "Homo sapiens 3-oxoacid CoA transferase"; Synthetic cos	24.1	20.3	26.5	46.3	5e-37	01.04.01 phosphate utilization
Pc14g0081	weak similarity to sphingoid long chain base kinases Lcb4 - Saccharomyces cerevisiae	1e-07	AB018110 Arabidopsis thaliana genomic DNA, chromosome 5, TAC cloneK19A	131.3	185.6	278.0	406.8	1e-121	01.05 C-compound and carbohydrate metabolism
Pc15g0021	strong similarity to glucan 1,3-beta-glucosidase Bgl2 - Saccharomyces cerevisiae	2e-25	AF359360, gene: "OrfC"; product: "glucosidase"; Fusarium sporotrichoides trit	1180.7	1237.5	2819.4	2262.1	1e-132	01 METABOLISM
Pc15g0151	strong similarity to brown 2 protein abr2 - Aspergillus fumigatus	1e-77	AF104823, gene: "abr2"; product: "brown 2"; Aspergillus fumigatus brown 2 (abi	181.4	167.6	444.7	397.8	8e-75	01.05.04 regulation of C-compound and carbohydrate utilization
Pc16g0011	strong similarity to hypothetical protein RTS-beta - Homo sapiens	1e-118	AE013861, gene: "y2598"; product: "hypothetical"; Yersinia pestis KIM section 2T	12.0	12.0	15.2	25.8	7e-64	01 METABOLISM
Pc16g0011	strong similarity to high affinity hexose transporter Hxt1 - Saccharomyces cerevisiae	3e-77	S69591, sugar transport protein STP1 - yeast (Saccharomyces cerevisiae)	12.0	12.0	15.0	30.0	6e-82	01.01.04 regulation of amino acid metabolism
Pc16g0034	strong similarity to alcohol dehydrogenase pan2 - Mus musculus	1e-19	BC009830, product: "retinol dehydrogenase 14 (all-trans and 9-cis)"; Homo sapi	29.9	44.3	65.9	85.8	8e-76	01.05.07 C-compound, carbohydrate transport
Pc16g0351	strong similarity to thiamine repressible gene tht1p - Schizosaccharomyces pombe	8e-17	T40521, hypothetical protein SPBC530.05 - fission yeast (Schizosaccharomy	19.2	12.0	21.7	43.8	6e-69	01.05 C-compound and carbohydrate metabolism
Pc16g0361	strong similarity to myo-inositol transporter 2 ltr2p - Schizosaccharomyces pombe	5e-94	T43400, myo-inositol transporter - fission yeast (Schizosaccharomyces pombe)	80.1	61.2	168.0	199.2	0.0	01.06.01 lipid, fatty-acid and isoprenoid biosynthesis
Pc16g0371	strong similarity to enoyl reductase of the lovastatin biosynthesis locV - Aspergillus terreus	2e-78	TOXD_CO TOXD PROTEIN.	32.6	25.8	71.1	87.4	1e-157	01.05 C-compound and carbohydrate metabolism
Pc16g0381	strong similarity to polyketide synthase PKS1 - Cochlicobolus heterostrophus	0.0	AY495597, gene: "PKS7"; product: "polyketide synthase"; Gibberella moniformis	56.8	31.4	84.8	93.7	6e-93	01.05 C-compound and carbohydrate metabolism
Pc16g0521	strong similarity to alpha-xylosidase Xyl5 - Sulfolobus solfataricus	0.0	G83936, hypothetical protein BH2055 [imported] - Bacillus halodurans (strain	24.0	29.8	86.4	88.4	9e-80	01.05.07 C-compound, carbohydrate transport
Pc16g0571	strong similarity to thymidine synthase protein like protein An03g6740 - Aspergillus niger	2e-21	AC31370, conserved hypothetical protein Au05000 [Imported] - Agrobacterium t	114.8	133.0	125.6	327.1	1e-11	01.04.07 phosphate transport
Pc16g0681	strong similarity to high affinity glucose transporter HGT11 - Kluyveromyces fragilis [putative se	1e-68	A31776, lacase permease - yeast (Kluyveromyces marxianus var. lactis)	26.4	12.0	18.9	29.3	2e-79	01.05.07 C-compound, carbohydrate transport
Pc16g0691	strong similarity to allantate permease Dals - Saccharomyces cerevisiae	3e-68	T41604, probable membrane transport protein - fission yeast (Schizosacchar	12.0	12.0	13.5	35.8	1e-102	01 METABOLISM
Pc16g0841	strong similarity to maltose transport protein Mal31 - Saccharomyces cerevisiae	4e-99	BX842681, gene: "B13N4.080"; product: "probable maltose permease"; Neuros	52.8	67.5	75.5	152.7	0.0	01.01.01 amino acid biosynthesis
Pc16g0961	strong similarity to NAD(P)H-dependent xylose reductase XYL1 - Pachysolen tannophilus	1e-155	AF219625, gene: "xyrA"; product: "D-xylose reductase"; Aspergillus niger D-xylo	14.7	16.6	33.1	64.8	5e-99	01.03 nucleotide metabolism
Pc16g1001	strong similarity to sulfite permease SubB - Penicillium chrysogenum	0.0	AF163974, gene: "subB"; product: "sulfate permease SubB"; Penicillium chrysog	472.9	577.1	926.6	1470.0	0.0	01 METABOLISM
Pc16g1051	strong similarity to adenosine deaminase Aah1 - Saccharomyces cerevisiae	1e-133	AF123480, gene: "nsdA"; product: "adenine deaminase"; Emmericella nidulans ad	14.3	21.5	32.3	41.3	9e-75	01.05.04 regulation of C-compound and carbohydrate utilization
Pc16g1121	strong similarity to copper amino oxidase AO-1 - Aspergillus niger	1e-163	BX64906, gene: "ao"; product: "copper amino oxidase 1, putative"; Aspergillu	12.0	12.0	17.1	27.4	3e-76	25.05.25 gametogenesis
Pc16g1121	strong similarity to hexose transporter ght2p - Schizosaccharomyces pombe	2e-93	NC13E11, gene: "13E11.140"; product: "probable sugar transporter"; Neurosp	12.7	12.0	41.3	42.2	1e-101	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc16g1131	strong similarity to novobiosin biosynthetic gene novR - Streptomyces spheroides	1e-69	AL646066, gene: "RSCT1767"; "RSO2954"; product: "CONSERVED HYPOTHET	221.1	414.4	430.5	967.3	0	
Pc16g1161	strong similarity to hypothetical sterigmatocystin biosynthesis p450 monooxygenase stcB - A	8e-71	AF169016, gene: "cypC"; product: "cytochrome P450 monooxygenase"; Aspergill	12.0	12.0	41.1	19.4	0	
Pc16g1161	strong similarity to arylsulfate sulfotransferase like protein An13g03160 - Aspergillus niger	2e-57	BX649067, product: "hypothetical protein"; Aspergillus fumigatus BAC pilot proje	38.0	33.7	54.0	77.8	5e-81	01.05 C-compound and carbohydrate metabolism

Pc180477	strong similarity to phenol 2-monooxygenase - Trichosporon belgelli	66-88	BX309810 gene: "G3C5.110"; product: "probable phenol 2-monooxygenase"; Ne	12.0	12.0	23.1	30.8	1e-179	01.01.01 amino acid biosynthesis
Pc180404	weak similarity to myo-inositol 2-dehydrogenase yJcS - Bacillus subtilis	1e-177	BX482620 gene: "B1T5E.410"; product: "conserved hypothetical protein"; Neur	20.6	25.5	28.3	52.1	1e-118	01.05.01 C-compound and carbohydrate utilization
Pc180133	strong similarity to saccharopine reductase LYV3 - Magnaporthe oryzae [putative sequencing]	1e-133	T40337 probable saccharopine dehydrogenase (NADP-, L-glutamate-forming)	80.4	121.8	168.8	267.4	1e-32	01.02 nitrogen and sulfur metabolism
Pc180156	strong similarity to glycolate oxidase GOX - Mus musculus	4e-78	AF140312 product: "glycolate oxidase"; Mus musculus glycolate oxidase mRNA	14.0	16.5	19.8	35.9	2e-82	01.01 amino acid metabolism
Pc180212	similarity to epoxide hydrolase - Rattus norvegicus	8e-54	AF27333 hypothetical protein af4221 [Imported] - Nostoc sp. (strain PCC 7120	29.1	34.9	39.8	104.8	3e-98	01.05 C-compound and carbohydrate metabolism
Pc180244	strong similarity to high affinity iron-binding protein Mup1 - Saccharomyces cerevisiae	1e-160	NC019430 gene: "HIF1"; product: "high-affinity iron-binding protein"; Schizosac	122.0	194.3	260.0	476.0	1e-117	01.01 amino acid metabolism
Pc180488	strong similarity to formaldehyde diismutase like protein AtG707160 - Aspergillus niger	5e-17	BX752599 product: "zinc-containing [hypothetical]; Rhodospirillum rubrum	12.0	19.8	43.8	48.1	2e-81	01.05.01 C-compound and carbohydrate utilization
Pc180498	strong similarity to alkane-inducible cytochrome P450 allele ALK1 - Yarrowia lipolytica	3e-87	AB010388 gene: "ALK1"; product: "ALK1"; Yarrowia lipolytica gene for ALK1, c	51.6	56.9	150.0	236.0	1e-132	01.03.19 nucleotide transport
Pc180555	similarity to 6-hydroxy-D-nicotinic oxide 6-HNDO - Arthrobaacter oxauidans	2e-19	AP003010 gene: "mlr6875"; product: "probable oxioreductases"; Mesorhizobiu	208.3	189.9	312.5	462.7	4e-70	01.05.04 regulation of C-compound and carbohydrate utilization
Pc200914	strong similarity to purine permease with broad specificity aspar - Aspergillus nidulans	0.0	A62843 purine permease, broad specificity - Emmericella nidulans	16.9	24.3	37.6	12.2	1e-62	01 METABOLISM
Pc200917	strong similarity to geranylgeranyl pyrophosphate synthase h3 - Neurospora crassa	1e-122	MG17_11_KLUGHSIAF-RANSPORC1 - Kluyveromyces fragilis	1e-122	1e-122	1e-122	1e-122	1e-122	01.02 nitrogen and sulfur metabolism
Pc200322	3-phosphoadenosine-5-phosphotransferase ParA - Penicillium chrysogenum	1e-180	AF576382 gene: "Para"; product: "3'-phosphoadenosine-5'-phosphotransfer red	297.6	354.1	487.6	825.8	4e-80	01.05.07 C-compound, carbohydrate transport
Pc200385	strong similarity to epoxide hydrolase hyf1 - Aspergillus niger	1e-140	ANI238455 gene: "hyf1"; product: "epoxide hydrolase"; Aspergillus niger hyf1 ge	64.2	95.1	198.8	236.3	1e-63	01.01.11.1 biosynthesis of the cysteine-aromatic group
Pc200403	strong similarity to quinate transport protein qutD - Aspergillus nidulans	1e-58	BX496055 gene: "qutD"; product: "quinat permease, putative"; Aspergillus fun	458.9	93.3	128.0	605.9	1e-98	01.01.04 regulation of amino acid metabolism
Pc200426	strong similarity to levodone reductase lrv - Corynebacterium aquificum	4e-30	AE012980 gene: "FabG"; product: "Dehydrogenases with different specificities	186.3	220.3	287.3	508.8	2e-60	01.01.11.1 biosynthesis of the cysteine-aromatic group
Pc200466	similarity to transcription factor ArgR - Saccharomyces cerevisiae	1e-43	S49827 regulatory protein ARG22 - yeast (Saccharomyces cerevisiae)	12.5	18.7	30.1	55.3	4e-81	01.05.07 C-compound, carbohydrate transport
Pc200498	strong similarity to 3-oxohexanoate decarboxylase Kds36.13 - Schizosaccharom	1e-100	AF166337 gene: "KDS36.13"; product: "3-oxohexanoate decarboxylase"; Schizo	135.3	157.7	181.4	314.4	1e-99	01.01.04 regulation of amino acid metabolism
Pc200772	strong similarity to mitochondrial phosphate transporter Grt - Glycine max	1e-117	TAC238533 gene: "ppc"; product: "putative mitochondrial phosphate carrier prote	153.3	254.5	851.0	647.4	1e-75	01.01.04 regulation of amino acid metabolism
Pc200934	similarity to hypothetical ureidoglycolate hydrolase SPAC1927.04 - Schizosaccharom	9e-19	SM2022 hypothetical ureidoglycolate hydrolase (EC 3.5.3.1) - yeast (Saccharomyc	19.7	31.1	50.2	61.2	1e-61	01.05.01 C-compound and carbohydrate utilization
Pc200966	similarity to transcription factor AroB - Saccharomyces cerevisiae	3e-23	SG69704 hypothetical protein YDR421+ve - yeast (Saccharomyces cerevisiae)	14.5	12.0	39.4	33.0	1e-90	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc200992	similarity to 6-hydroxy-D-nicotinic oxide 6-HNDO - Arthrobaacter oxauidans	6e-54	T48777 6-HYDROXY-D-NICOTINIC OXIDASE related protein [imported] - Ne	13.4	12.0	67.8	14.7	1e-133	01.06.01 lipid, fatty-acid and isoprenoid biosynthesis
Pc200998	strong similarity to geranylgeranyl pyrophosphate synthase h3 - Neurospora crassa	0.0	MC127612 gene: "cgs3"; product: "geranylgeranyl pyrophosphate synthase"; M	8.6	11.0	13.8	16.9	1e-69	01 METABOLISM
Pc201025	strong similarity to sterol delta14,15-reductase E3 - Neurospora crassa	0.0	H507326 gene: "STR2"; product: "sterol delta14,15-reductase"; Neurospora	153.4	186.9	113.9	218.1	1e-108	01.01 amino acid metabolism
Pc201047	strong similarity to delta3-cis-delta2-trans-enoyl-CoA isomerase Ed1 - Saccharomyc	7e-41	MS2399 probable membrane protein YLR284c; - yeast (Saccharomyces cerevi	48.2	118.2	153.3	300.8	1e-135	01 METABOLISM
Pc201043	strong similarity to biotin synthesis enzyme like protein AN1501990 - Aspergillus niger	1e-121	AP003859 gene: "UJ1033_B09.17"; product: "putative adenosinemethionine-8-an	226.3	193.1	372.1	568.6	0	01.05.01 C-compound and carbohydrate utilization
Pc201356	strong similarity to very long-chain fatty acyl-CoA synthase Fat1 - Saccharomyc	1e-163	ACH507635 gene: "celD1"; product: "isopenicillin N-CoA synthetase"; Acetromu	54.7	25.9	45.7	143.7	1e-108	01.02 secondary metabolism
Pc201351	strong similarity to methylcrotonyl carboxylase hmaA - Aspergillus nidulans	0.0	ANI249111 gene: "hmaA"; product: "methylcrotonyl synthase"; Aspergillus nidul	254.2	229.7	319.8	460.0	5e-81	01.05 C-compound and carbohydrate metabolism
Pc201356	strong similarity to very long-chain fatty acyl-CoA synthase Fat1 - Saccharomyc	1e-163	ACH507635 gene: "celD1"; product: "isopenicillin N-CoA synthetase"; Acetromu	54.7	25.9	45.7	143.7	1e-108	01.02 secondary metabolism
Pc201405	similarity to fructoseamine 3-kinase FN3K - Homo sapiens	3e-27	NCB9814_4 gene: "BC3E4.080"; product: "conserved hypothetical protein"; Neur	19.7	13.5	28.4	37.1	1e-102	01.10.10 amino acid degradation (catabolism)
Pc201488	strong similarity to general amino acid permease Gap1 - Saccharomyc	1e-77	UFA30825 gene: "AAT1"; product: "amino acid transporter"; Uromyces fabae A	12.0	12.0	12.0	40.3	0	01.06.01 lipid, fatty-acid and isoprenoid biosynthesis
Pc201515	similarity to fructosyl amine oxygen oxidoreductase - Aspergillus fumigatus	5e-23	T40295 fructosyl amine - fission yeast (Schizosaccharomyc	55.1	103.0	116.2	217.3	9e-75	01.05.07 C-compound, carbohydrate transport
Pc210098	strong similarity to lovastatin diphosphate synthase lovF - Aspergillus terreus	0.0	AY495602 gene: "PKS12"; product: "polyketide synthase"; Gibberella moniliform	81.0	16.6	138.1	116.3	1e-148	01.05.01.01 C-compound, carbohydrate catabolism
Pc210172	strong similarity to quinate transport protein qutD - Aspergillus nidulans	2e-82	S08498 quinate transport protein - Emmericella nidulans	48.8	36.5	73.7	17.1	1e-53	01.01.11.1 biosynthesis of the cysteine-aromatic group
Pc210235	strong similarity to hecA-like protein hecA - Aspergillus nidulans	0.0	MC392833 gene: "hecA"; product: "hecA-like protein"; Aspergillus nidulans	33.9	164.1	164.1	164.1	1e-100	01.01.01 amino acid biosynthesis
Pc210305	strong similarity to meso-2,3-butanediol dehydrogenase like protein AN1405430 - Aspergillus	3e-42	F83629 probable short chain dehydrogenase PA0117 [Imported] - Pseudomon	35.7	42.8	58.6	94.4	0	0
Pc210355	strong similarity to tryptophan synthase Trp5 - Saccharomyc	0.0	AF070933 gene: "trpB"; product: "bifunctional tryptophan synthase TRPB"; Asp	207.2	197.1	395.3	404.4	1e-50	01.05.07 C-compound, carbohydrate transport
Pc210441	similarity to cytosolic deaminase codA - Escherichia coli	6e-26	AE016948 product: "N-acyl-D-amino-acid deacylase family protein"; Enterococc	38.1	30.5	58.1	80.5	8e-80	01.01.04 regulation of amino acid metabolism
Pc210458	strong similarity to alanine permease - Saccharomyc	9e-89	T41345 probable alanine permease - fission yeast (Schizosaccharomyc	30.0	27.4	50.2	104.0	4e-46	01.01.11.1 biosynthesis of the cysteine-aromatic group
Pc210464	weak similarity to putative amino acid transporter - Neurospora crassa	0.0	AF218847 gene: "PRF"; product: "putative amino acid transporter"; Neurospora	19.7	13.5	28.4	37.1	1e-102	01.05.07 C-compound, carbohydrate transport
Pc210481	strong similarity to developmental protein factor csfA - Pyricularia oryzae	2e-32	AG03988 probable short-chain dehydrogenase [Imported] - Yersinia pestis (stra	12.0	14.4	14.1	32.3	1e-14	01.05.01 resistant proteins
Pc210834	strong similarity to quinate transport protein qutD - Aspergillus nidulans [putative sequencing]	2e-67	S08498 quinate transport protein - Emmericella nidulans	46.2	28.1	56.1	102.3	2e-84	01.05.07 C-compound, carbohydrate transport
Pc210961	strong similarity to esterase A gene EstA - Streptomyces chrysomallus	1e-101	AY052630 gene: "cddB"; product: "lactone dehydratase"; Rhodococcus ruber strai	116.2	57.8	130.3	202.8	3e-92	01.01.11.1 biosynthesis of the cysteine-aromatic group
Pc210968	strong similarity to quinate transport protein qutD - Aspergillus nidulans	4e-86	G31727 quinate transport protein - Neurospora crassa (tentative sequencing)	12.0	12.0	13.7	32.5	4e-95	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc210978	strong similarity to fructosyl amine oxygen oxidoreductase - Aspergillus fumigatus	0.0	AF035700 product: "fructosyl amine oxygen oxidoreductase"; Aspergillus fumigi	12.0	12.0	30.7	43.3	0.0	01.05 C-compound and carbohydrate metabolism
Pc211019	strong similarity to alanine permease - Saccharomyc	1e-118	AY102652 gene: "TR4"; product: "cytosolic alanine permease"; Neurospora	12.0	12.0	12.0	12.0	1e-118	01.05.01 amino acid metabolism
Pc211008	strong similarity to 1,4-Ala-branch branching enzyme Glc3 - Saccharomyc	0.0	AB077230 gene: "gbeA"; product: "glycogen branching enzyme"; Aspergillus or	129.6	155.9	370.4	388.0	1e-73	01.01.07 biosynthesis of vitamins, cofactors, and prosthetic groups
Pc211025	strong similarity to salicylate hydroxylase nahG - Pseudomonas putida	4e-53	BX640431 product: "putative monooxygenase"; Bordetella parapertussis strain	78.1	64.8	106.7	131.0	3e-46	01.05.01 C-compound and carbohydrate utilization
Pc211025	strong similarity to pyridoxamine-phosphate oxidase Pdc3 - Saccharomyc	2e-36	S41301 pyridoxamine-phosphate oxidase (EC 1.4.3.5) - yeast (Saccharomyc	19.5	26.5	63.0	53.9	2e-51	01.04.07 phosphate transport
Pc211045	strong similarity to carbonic anhydrase pcal1 - Porphyridium purpurum	8e-55	NI012245 gene: "pcal1"; product: "Carbonic anhydrase"; Leptospira interrogans s	37.6	33.8	133.6	96.7	0	0
Pc211059	strong similarity to alanine permease - Saccharomyc	1e-118	T41345 probable alanine permease - fission yeast (Schizosaccharomyc	30.0	27.4	50.2	104.0	4e-46	01.05.01 amino acid metabolism
Pc211165	weak similarity to hypothetical polycarboxylate synthase CP51 - Filobasidiella neoforma [ju	1e-84	BX482620 gene: "B1C55.060"; product: "putative protein"; Neurospora crassa	12.0	12.0	12.0	12.0	1e-84	01.06.04 breakdown of lipids, fatty acids and isoprenoids
Pc211175	strong similarity to 3-dehydroshikimate dehydratase ga-4 - Neurospora crassa	2e-57	BX496055 gene: "qutC"; product: "3-dehydroshikimate dehydratase, putative"; f	12.0	12.0	12.0	32.1	1e-164	01.05.01.01 C-compound, carbohydrate catabolism
Pc211211	strong similarity to triacylglycerol lipase Tgl2 - Saccharomyc	4e-40	SC2TGL2_1 gene: "TGL2"; product: "triglyceride lipase"; S. cerevisiae TGL2 ge	56.5	53.5	75.4	121.7	5e-67	01.03 nucleotide metabolism
Pc211367	strong similarity to enzyme with sugar transferase activity like protein AN0110930 - Aspergill	0.0	AB057788 gene: "agbB"; product: "alpha-glucuronidase"; Aspergillus nidulans g	881.8	1205.8	3599.5	3331.6	3e-23	01.01.04 regulation of amino acid metabolism
Pc211498	strong similarity to methionine adenosyltransferase regulatory beta subunit like protein C203-967	3e-97	MS101044 gene: "B1014.170"; product: "related to methionine adenosyltransfer	36.9	44.8	84.3	110.8	4e-90	01.04 phosphate metabolism
Pc211501	strong similarity to transaminase - Saccharomyc	1e-100	NC019430 gene: "HIF1"; product: "high-affinity iron-binding protein"; Schizosac	122.0	194.3	260.0	476.0	1e-117	01.01.04 lipid, fatty-acid and isoprenoid metabolism
Pc211525	strong similarity to phosphate-repressible phosphate permease pho4 - Neurospora crassa	2e-85	JQ01116 phosphate-repressible phosphate permease - Neurospora crassa	37.6	36.8	77.2	45.3	1e-143	01.04.01 phosphate utilization
Pc211545	strong similarity to cytochrome P450 monooxygenase P450 - Gibberella fujikuri	8e-98	GFP4450_1 gene: "P450"; product: "cytochrome P450 monooxygenase"; Gibber	344.1	281.2	1420.2	617.7	1e-112	01.02.01.09.9 other catabolism of nitrogenous compounds
Pc211591	strong similarity to cytosolic exopolyphosphatase Ppx1 - Saccharomyc	4e-24	T38544 probable exopolyphosphatase - fission yeast (Schizosaccharomyc	31.2	29.5	54.8	83.3	1e-124	01.01.15.03 biosynthesis of leucine
Pc211603	strong similarity to dimethylamine monooxygenase like protein AN0605690 - Aspergillus nigr	3e-94	NC933111 gene: "93G5.11.170"; product: "related to flavin-containing monooxyge	21.5	23.5	38.0	62.1	0.0	01 METABOLISM
Pc211755	strong similarity to acyl-CoA dehydrogenase like protein AN1701150 - Aspergillus niger	0.0	NC981844 gene: "B1014.130"; product: "conserved hypothetical protein"; Neur	529.8	553.1	839.7	1248.7	1e-120	01.05.01 C-compound and carbohydrate utilization
Pc211775	strong similarity to glutamine synthetase glnA - Aspergillus nidulans	1e-81	SO36721_1 gene: "glnA"; product: "glutamine synthetase"; Pichia pastoris	18.0	24.0	34.4	36.9	1e-77	01.01 amino acid metabolism
Pc211781	strong similarity to flavonoylchome b2 L-lactate dehydrogenase CYB2 - Pichia anomala	1e-62	AB020451 gene: "agaA"; product: "alpha-glucuronidase"; Aspergillus niger aga	12.0	12.0	12.0	40.3	0.0	01.01.01 amino acid biosynthesis
Pc211815	strong similarity to alpha-glucuronidase agaA - Aspergillus niger	0.0	T38544 probable exopolyphosphatase - fission yeast (Schizosaccharomyc	31.2	29.5	54.8	83.3	1e-124	01.01.15.03 biosynthesis of leucine
Pc211875	strong similarity to hypothetical protein AN02030790 - Aspergillus niger	1e-139	T39195 phosphate adeno acid permease - fission yeast (Schizosaccharomyc	90.8	123.2	286.5	367.5	1e-36	99 UNCLASSIFIED PROTEINS
Pc211938	strong similarity to salicylate hydroxylase nahG - Penicillium chrysogenum	0.0	AF169375 gene: "nahG"; product: "salicylate hydroxylase"; Aspergillus niger	34.2	12.0	35.5	51.4	1e-80	01.01.99 other amino acid metabolism activities
Pc212004	strong similarity to alanine permease - Saccharomyc	1e-118	T41345 probable alanine permease - fission yeast (Schizosaccharomyc	30.0	27.4	50.2	104.0	4e-46	01.05.01 amino acid metabolism
Pc212138	isoprenoid N synthase ps PCBC - Penicillium chrysogenum	0.0	S04441 isopenicillin N synthase (EC 1.14.11.3) [pe] [similarity] - Penicillium ch	1312.0	2381.9	5107.0	5111.8	1e-158	13 REGULATION OF / INTERACTION WITH CELLULAR ENVIRONMENT
Pc212135	alpha-aminoacid-pyruvyl-cystenyl-valine synthetase pcbAB acvA - Penicillium chrysogenum [putat	0.0	YGPLV8 alpha-aminoacid-pyruvyl-cystenyl-valine synthetase (EC 6.-.-.-) - Penicilliu	1098.0	2060.0	4345.2	4557.4	1e-66	01.01.11.1 biosynthesis of the cysteine-aromatic group
Pc212154	strong similarity to lipase lip1 - Geotrichum candidum	1e-103	NCB23231_1 gene: "B23G1.090"; product: "related to cholinesterase"; Neurosp	12.0	12.4	27.8	53.8	1e-134	01 METABOLISM
Pc212181	strong similarity to levodone reductase lrv - Corynebacterium aquificum	1e-110	AF11A2H2_2 gene: "A11A2H2.17c"; product: "putative 3-oxoacyl-acyl-carboxylate	79.3	82.0	154.4	241.3	1e-90	01 METABOLISM
Pc212245	strong similarity to p-nitrobenzyl esterase pbp1 - Bacillus subtilis	7e-33	YU1249751 gene: "fbp3"; product: "carboxylesterase/lipase type B"; Yarrowia lipot	12.0	12.0	42.4	50.8	1e-104	01 METABOLISM
Pc212255	strong similarity to hydroxymethylglutaryl-CoA synthase - Saccharomyc	1e-100	CA23881_2 product: "Porphyridium purpurum hydroxymethylglutaryl-CoA synth	12.0	12.0	12.0	12.0	1e-100	01.05.01 amino acid metabolism and sulfur utilization
Pc212265	strong similarity to aldehydehydroxylase ALR - Rattus norvegicus	6e-73	AY207463 gene: "AlrA1"; product: "aldehyde reductase"; Mus musculus aldH	12.0	12.0	12.0	41.8	2e-93	01 METABOLISM
Pc212355	strong similarity to cyanamide hydratase - Myrothecium verrucaria	6e-66	S56194 hypothetical protein YFL061+ve - yeast (Saccharomyc	193.0	191.7	111.0	409.3	6e-84	01 METABOLISM
Pc212365	strong similarity to D-lactate dehydrogenase ldhA - Escherichia coli	1e-127	BX482620 gene: "90C4.110"; product: "related to D-lactate dehydrogenase"; Ne	82.9	87.7	155.3	108.9	3e-64	01.01.11.1 biosynthesis of the cysteine-aromatic group
Pc220021	strong similarity to protein with seroselective transaminase activity like protein AN0408630	3e-66	ML067221 gene: "msd247"; product: "PUTATIVE BRANCHED-CHAIN AMINO, I	53.5	29.5	52.8	105.3	1e-79	01.05.07 C-compound, carbohydrate transport
Pc220032	strong similarity to short-chain alcohol dehydrogenase adhA - Pseudomonas parasitica	8e-15	AB067878 gene: "bacC"; product: "BacC"; Bacillus subtilis strain A13 bacilysin	85.6	84.4	16.2	273.0	1e-168	01 METABOLISM
Pc220055	strong similarity to carnitine acyl-CoA transferase - Saccharomyc	1e-124	CA23881_2 product: "Porphyridium purpurum carnitine acyl-CoA transferase	12.0	12.0	12.0	12.0	1e-124	01.05.01 amino acid metabolism
Pc220096	strong similarity to acetoacetyl-CoA synthase acsA - Sinorhizobium meliloti	0.0	NC151111 gene: "15E11.130"; product: "related to acetoacetyl-CoA synthetase"	35.6	71.6	112.8	106.1	1e-126	01.01 amino acid metabolism
Pc220091	strong similarity to glutamate decarboxylase Gad1 - Arabidopsis thaliana	0.0	JC7915 glutamate decarboxylase (EC 4.1.1.15) - Aspergillus oryzae	54.2	61.9	161.9	205.6	3e-70	01 METABOLISM
Pc220091	strong similarity to glutamate decarboxylase Gad1 - Arabidopsis thaliana	0.0	JC7915 glutamate decarboxylase (EC 4.1.1.15) - Aspergillus oryzae	54.2	61.9	161.9	205.6	3e-70	01.01.10 amino acid degradation (catabolism)
Pc220107	strong similarity to nitrilase NIT1 - Arabidopsis thaliana	1e-144	AB087872 gene: "NIT1"; product: "hypothetical nitrilase-like protein"; Aspergillu						

Pc22g090: strong similarity to 3-oxoacyl-[acyl-carrier-protein] synthase like protein An02g14220 - Asperg	1e-143	AF021234	gene: "cem-1"; product: "3-oxoacyl-[acyl-carrier-protein]-synthase"; N	53.6	51.6	84.1	111.2	6e-64	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc22g094: strong similarity to aryl-alcohol oxidase precursor aao - Pleurotus eryngii	1e-160	CNS0771X	DNA centromeric region; produced from BAC DP15B03, DP38F06 of	12.0	12.0	13.1	33.0	3e-66	01.06.01 lipid, fatty-acid and isoprenoid biosynthesis
Pc22g094: strong similarity to 2-keto-3-deoxyglucose oxidoreductase kdD - Erwinia chrysanthemi	1e-40	TACD3.9	gene: "Ta0747"; product: "glucose 1-dehydrogenase related protein";	63.3	35.6	49.4	99.9	6e-55	01.02.01 nitrogen and sulfur utilization
Pc22g095: strong similarity to cyclopropane fatty-acyl-phospholipid like protein An02g0530 - Aspergillu	1e-36	AP005038	gene: "cfa"; product: "putative cyclopropane fatty acid synthase";	23.5	23.5	71.2	131	2e-72	01.06.10 regulation of lipid, fatty-acid and isoprenoid metabolism
Pc22g114: strong similarity to 2-nitropropane dioxygenase ncd-2 - Neurospora crassa	8e-55	T46693	probable 2-nitropropane dioxygenase (EC 1.13.1.32) precursor [mp	187.5	248.9	50.9	622.0	0.0	01.04 phosphate metabolism
Pc22g120: strong similarity to protein involved in fatty acid regulation like protein An04g07120 - Aspergill	3e-48	AE017009	product: "Nitroreductase family protein"; Bacillus cereus ATCC 1457	51.2	65.8	78.0	161.8	9e-73	01.01.04 regulation of amino acid metabolism
Pc22g123: strong similarity to extracellular alpha-glucosidase agU - Aspergillus niger	0.0	JC4217	alpha-glucosidase (EC 3.2.1.20) - Aspergillus oryzae	70.2	70.3	297.0	214.2	3e-55	01.05.04 regulation of C-compound and carbohydrate utilization
Pc22g123: strong similarity to transcription regulator of maltose utilization amyR - Aspergillus oryzae	0.0	AF155808	gene: "amyR"; product: "unknown"; Aspergillus niger (amyR) gene,	121.5	83.0	353.7	458.8	1e-76	01.05 C-compound and carbohydrate metabolism
Pc22g127: strong similarity to hypothetical sugar transporter encoded by DRA0271 - Deinococcus radio	7e-96	SPAP81E	gene: "SPAP81E7.08c"; S.pombe chromosome I BAC pB1E7;	29.9	25.3	36.4	59.8	1e-129	01.01 amino acid metabolism
Pc22g133: strong similarity to alcohol dehydrogenase like protein An02g0200 - Aspergillus niger	1e-41	AE017006	product: "Alcohol dehydrogenase"; Bacillus cereus ATCC 14579 ser	288.1	301.5	520.1	801.7	4e-78	01.05.07 C-compound, carbohydrate transport
Pc22g136: strong similarity to 2,2-dialkylglycine decarboxylase structural protein dgda - Pseudomonas	c1e-171	AY039312	gene: "DGD1"; product: "dialkylglycine decarboxylase"; Mycoplasma	12.0	12.0	20.6	37.8	1e-130	01 METABOLISM
Pc22g145: strong similarity to maltose transport protein Mal31 - Saccharomyces cerevisiae	1e-96	BX842681	gene: "B13N4.080"; product: "probable maltose permease"; Neuros	76.4	60.4	94.5	159.0	7e-94	01.05.01 C-compound and carbohydrate utilization
Pc22g148: phenylacetyl-CoA ligase pcdA - Penicillium chrysogenum	0.0	PCPHCOA	gene: "pcdA"; product: "phenylacetyl-CoA ligase"; Penicillium chrys	108.5	140.0	222.5	381.6	1e-100	01.05.01 C-compound and carbohydrate utilization
Pc22g153: strong similarity to alpha-glucan synthase mck1p - Schizosaccharomyces pombe	0.0	T43731	cell wall alpha-glucan synthase (EC 2.4.1.-) mck1 - fission yeast	69.5	35.9	101.0	149.1	1e-82	01.07.01 biosynthesis of vitamins, cofactors, and prosthetic groups
Pc22g165: strong similarity to isomalt alcohol oxidase mreA - Aspergillus oryzae	0.0	AB078977	gene: "mreB"; product: "probable alcohol oxidase"; Aspergillus oryza	118.0	112.8	182.5	242.2	1e-127	01 METABOLISM
Pc22g193: strong similarity to biotin-precursor ligase Bp11 - Saccharomyces cerevisiae [putative sequ	1e-111	S64646	gene: "SPAP81E7.08c"; S.pombe chromosome I BAC pB1E7;	82.2	95.5	111.3	174.1	1e-75	01 METABOLISM
Pc22g202: strong similarity to 4-coumarate-CoA ligase Acl - Arabidopsis thaliana	2e-94	SC093931	gene: "SC04383"; "Acl.2"; "SCD10.15"; product: "4-coumarate-CoA	11.18	173.9	186.3	458.8	0.0	01.05 C-compound and carbohydrate metabolism
Pc22g203: strong similarity to carnitine racemase like protein An03g03550 - Aspergillus niger	6e-40	AP005945	gene: "fadB"; product: "enoyl-CoA hydratase"; Bradyrhizobium japoni	12.0	12.0	12.0	43.3	1e-126	01.01.01 amino acid biosynthesis
Pc22g221: strong similarity to 4-alpha-glucanotransferase amylo-1,6-glucosidase Gdb1 - Saccharomy	0.0	S59841	4-alpha-glucanotransferase / amylo-1,6-glucosidase homolog YPR18	126.0	175.2	750.7	460.1	8e-61	01.04.07 phosphate transport
Pc22g225: strong similarity to branched-chain amino acid aminotransferase bcaT - Lactococcus lactis	5e-81	AF157629	gene: "TOXP"; product: "putative branched-chain amino acid aminotr	140.9	173.4	442.5	370.7	1e-157	01.05 C-compound and carbohydrate metabolism
Pc22g234: strong similarity to allantoate permease Da5 - Saccharomyces cerevisiae	3e-84	T41604	probable membrane transport protein - fission yeast (Schizosacchar	12.0	12.0	12.0	46.1	0	0
Pc22g233: strong similarity to aldehyde dehydrogenase adh3 - Homo sapiens	8e-88	AF164552	gene: "TG0H4.133"; product: "Hypothetical protein TG0H4.13b"; Caen	188.1	110.5	162.9	279.8	4e-79	01.01.01.01 assimilation of ammonia, biosynthesis of the glutamate group
Pc22g237: weak similarity to fructosyl amine oxygen oxidoreductase - Aspergillus fumigatus	1e-04	AF162830	product: "fructosyl amine oxygen oxidoreductase"; Aspergillus fumig	12.4	15.1	32.4	52.1	1e-45	01.05 C-compound and carbohydrate metabolism
Pc22g237: strong similarity to pyridine-5-carboxylate reductase PSOR - Zalerion arboricola	2e-55	JC4830	pyrroline-5-carboxylate reductase (EC 1.5.1.2) - Zalerion arboricola	12.0	14.4	25.3	37.1	1e-124	01.05.01 C-compound and carbohydrate utilization
Pc22g239: strong similarity to hypothetical phosphoglycerate mutase YKR043c - Saccharomyces cerevi	2e-45	S38115	hypothetical protein YKR043c - yeast (Saccharomyces cerevisiae)	44.0	36.1	101.0	123.7	4e-83	01.05.01 C-compound and carbohydrate utilization
Pc22g241: strong similarity to L-lactate 2-monooxygenase LA2M - Mycobacterium smegmati	1e-106	SPAPB1A	gene: "SPAPB1A11.03"; S.pombe chromosome I BAC pB1A11.	106.8	73.2	219.3	451.4	1e-116	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc22g248: strong similarity to 6-hydroxy-d-nicotine oxidase 6-HNDO - Anthrobacter oxidans	4e-51	B59597	probable oxidoreductase, oxygen dependent, FAD-dependent protein	83.1	109.4	160.4	230.3	3e-95	01 METABOLISM
Pc22g251: strong similarity to acyl CoA dehydrogenase adhB - Escherichia coli	4e-81	NC2E4_2	gene: "2E4.0207"; product: "conserved hypothetical protein"; Neuros	46.2	17.5	66.9	198.8	5e-61	01.01.10.01.03 degradation of arginine
Pc22g251: strong similarity to chitinase 1 precursor c1 - Coccidioides immitis	6e-97	AF529207	gene: "chB"; product: "endochitinase"; Botryotinia fuckeliana endoch	156.3	165.3	969.6	1466.8	1e-121	01.04 metabolism of energy reserves (e.g. glycogen, trehalose)
Pc12g146: strong similarity to flavohemoglobin Flp - Alcaligenes eutrophus	1e-104	AB016807	product: "flavohemoglobin"; Fusarium oxysporum mRNA for flavohem	148.3	150.1	243.0	430.1	7e-42	01.01.01.01.02.01 biosynthesis of proline
Pc16g042: strong similarity to trehalose metabolism factor Pmmt - Saccharomyces cerevisiae	2e-32	T18237	conserved hypothetical protein - yeast (Candida albicans)	31.5	30.8	34.8	64.8	1e-109	01 METABOLISM
Pc20g020: strong similarity to 3-oxoadipate enol-lactone hydrolase catD - Pseudomonas sp.	1e-113	NCB1383	gene: "B1383.090"; product: "related to 3-OXOADIPATE ENOL-LAC	12.0	12.0	20.2	49.2	1e-100	01.03 nucleotide metabolism
Pc22g036: strong similarity to isocitrate lyase acd - Aspergillus nidulans	0.0	AY442291	Aspergillus fumigatus isocitrate lyase (icl) gene, complete cds.	276.0	155.8	456.4	509.8	1e-122	03.01.05 DNA recombination and DNA repair
Pc22g141: similarity to transaldolase talB - Synchocystis sp.	4e-25	AY144980	product: "TAL1"; Saccharomyces castellii clone Contig206 TAL1 g	34.7	41.9	59.7	130.1	3e-26	01.05.01.01 C-compound, carbohydrate catabolism
Pc13g05: similarity to hellicase P411 - Saccharomyces cerevisiae	5e-15	T47241	RIM4P411 helicase homolog - fission yeast (Schizosaccharomyces	37.8	22.9	50.7	58.3	1e-73	01.04 phosphate metabolism
Pc13g065: similarity to adaptive response regulator protein Ada - Escherichia coli	5e-19	F91016	OB-methylguanine-DNA methyltransferase [Imported] - Escherichia c	12.5	12.0	19.7	44.5	1e-120	01.05.01 DNA recombination and DNA repair
Pc16g138: similarity to ds1-suppressing protein kinase dsk1p - Schizosaccharomyces pombe	3e-12	S28282	hypothetical protein B0464.5 - Caenorhabditis elegans	55.7	39.2	86.9	83.8	0.0	03.05.01 DNA repair
Pc21g119: strong similarity to protein DLH1 - Candida albicans	1e-116	A38214	meiosis-specific recombination protein DMC1 [validated] - yeast (Sac	119.2	89.8	186.2	240.2	8e-68	01.04 phosphate metabolism
Pc22g225: strong similarity to DNA photolyase phr1 - Trichoderma harzianum	0.0	TA000996	gene: "phr1"; product: "DNA photolyase"; Trichoderma harzianum ph	14.7	12.0	43.0	29.8	4e-21	01.05.04 regulation of C-compound and carbohydrate utilization
Pc06g002: weak similarity to SR protein kinase SkY1 - Saccharomyces cerevisiae	4e-29	A47726	ds1-suppressing protein kinase dsk1 - fission yeast (Schizosacchar	26.7	12.0	38.6	34.1	6e-75	04 TRANSCRIPTION
Pc06g018: weak similarity to transcription activator 1 f yinase pathway Lys14 - Saccharomyces cerevi	1e-31	CNS09545	DNA centromeric region; sequence from BAC DP28B06, DP34F04, I	27.1	15.0	31.1	32.8	9e-38	01.01.04 regulation of amino acid metabolism
Pc06g020: similarity to developmental control protein like protein An09g01040 - Aspergillus niger	2e-06	AE003420	gene: "EG-BACR4217.11"; product: "CG11398-PA"; Drosophila mel	108.3	80.1	228.5	221.9	5e-79	01.06.01.07.11 tetracyclic and pentacyclic terpenes (cholesterol, steroids and hopanoids) biosynthesis
Pc13g065: strong similarity to root-specific homeotic protein like protein An14g00520 - Aspergillus niger	3e-15	BX842620	gene: "B11E5.180"; product: "hypothetical protein"; Neurospora cr	12.0	12.0	28.2	41.1	0	0
Pc13g094: strong similarity to transcription factor like protein An02g06940 - Aspergillus niger	7e-15	BC055035	gene: "Tcfq2p3"; product: "Tcfq2p3 protein"; Mus musculus transci	112.1	101.6	251.6	238.0	1e-90	01.01.04 regulation of amino acid metabolism
Pc13g098: weak similarity to cercosporin resistance protein crtG1 - Cercospora nicotianae	0	0	0	12.0	12.0	19.7	43.0	1e-107	01.01.04 regulation of amino acid metabolism
Pc13g144: strong similarity to cutinase transcription factor 1 CTF1-alpha - Fusarium solani	3e-29	CT1E_FUS	CUTINASE TRANSCRIPTION FACTOR 1 ALPHA	12.0	13.3	13.1	29.3	3e-21	01.01.04 regulation of amino acid metabolism
Pc4g000: weak similarity to hypothetical transcription regulator protein - Saccharomyces pombe	4e-24	T40521	hypothetical protein SPBC530.05 - fission yeast (Schizosaccharmy	65.6	132.7	453.2	348.2	7e-31	01.05.04 regulation of C-compound and carbohydrate utilization
Pc4g008: similarity to transcription activator of lysine pathway Lys14 - Saccharomyces cerevisiae	1e-06	T41710	hypothetical fungal Zv02Cys(6) a-gal - fission yeast - fission yeast (S	14.7	11.0	30.8	15.7	1e-67	04.05.01.04 transcriptional control
Pc4g013: similarity to transcription factor Upc2 - Saccharomyces cerevisiae	6e-08	S61580	probable membrane protein YDR213w - yeast (Saccharomyces cerev	94.7	144.9	235.0	320.1	0	0
Pc15g012: similarity to transcription regulator 2 of pal operon palB - Bacillus subtilis	2e-26	AP002994	gene: "mil0013"; product: "transcriptional regulator"; Mesorhizobium	88.3	83.5	174.2	169.7	1e-87	01.01.04 regulation of amino acid metabolism
Pc16g002: similarity to cutinase transcription factor alpha CTF1a - Fusarium solani	2e-14	NCB11N2	gene: "B11N2.030"; product: "probable CUTINASE TRANSCRIPTIC	12.0	12.0	16.2	40.3	2e-81	01.01.04 regulation of amino acid metabolism
Pc16g010: similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	4e-43	BX908809	gene: "29E8.250"; product: "conserved hypothetical protein"; Neuros	28.5	25.4	38.9	52.8	3e-80	04.05.01.04 transcriptional control
Pc16g091: strong similarity to nitrogen assimilation regulatory protein nit-4 - Neurospora crassa	3e-35	A41696	regulatory protein nit-4 - Neurospora crassa	18.1	19.6	66.0	62.8	9e-50	01.01.04 regulation of amino acid metabolism
Pc16g091: similarity to zinc-finger transcription factor andA - Aspergillus nidulans	5e-11	AY027529	product: "Ste12-like transcription factor"; Neurospora crassa Ste12-4	27.4	30.5	80.2	101.1	1e-31	01.01.04 regulation of amino acid metabolism
Pc18g016: weak similarity to protein PRB1 - Lentinus edodes	0	0	0	40.1	34.8	57.9	77.2	2e-18	01.01.04 regulation of amino acid metabolism
Pc18g031: weak similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces px	1e-08	T37604	probable transcription activator - fission yeast (Schizosaccharomyces	20.3	30.1	44.9	82.4	4e-37	01.01.04 regulation of amino acid metabolism
Pc20g017: similarity to transcription activator of lysine pathway Lys14 - Saccharomyces cerevisiae	9e-04	S61587	transcription activator LYS14 - yeast (Saccharomyces cerevisiae)	24.3	18.2	37.9	39.3	1e-103	01.03.13 regulation of nucleotide metabolism
Pc20g065: weak similarity to zinc-finger transcription factor andA - Aspergillus nidulans	8e-64	AJ582911	Aspergillus fumigatus negA gene for transcription factor RegA,	42.1	45.2	64.6	121.8	1e-95	01.05.04 regulation of C-compound and carbohydrate utilization
Pc20g065: similarity to hypothetical transcription regulator SPBC530.05 - Schizosaccharomyces pombe	6e-15	T40521	hypothetical protein SPBC530.05 - fission yeast (Schizosaccharmy	13.2	12.0	14.1	24.1	2e-97	01.01.04 regulation of amino acid metabolism
Pc20g074: similarity to protein SribB - Saccharomyces cerevisiae	2e-24	S74293	SRB9 protein - yeast (Saccharomyces cerevisiae)	10.0	12.0	20.4	27.1	6e-90	01.01.04 regulation of amino acid metabolism
Pc20g121: weak similarity to transcription activator Pu3 - Saccharomyces cerevisiae	2e-35	T38690	probable regulatory protein - fission yeast (Schizosaccharomyces po	12.0	12.0	13.2	33.3	0	0
Pc20g148: strong similarity to hypothetical transcriptional regulator SPCC417.09c - Schizosaccharmyc	1e-130	ANAJ3459	gene: "pmaA"; product: "PmaA protein"; Aspergillus nidulans pmd, prm	19.5	27.1	62.1	78.8	5e-75	01.04 phosphate metabolism
Pc20g150: similarity to aflatoxin biosynthesis regulator alR - Aspergillus flavus	6e-05	AF441421	gene: "alR"; product: "AflR"; Aspergillus nomius isolate BN013P A	32.3	26.4	47.1	70.4	0	0
Pc21g005: similarity to SR protein kinase SkY1 - Saccharomyces cerevisiae	7e-27	CA8553	probable protein kinase [Imported] - Arabidopsis thaliana	17.6	12.0	14.4	28.8	8e-58	01.01.04 regulation of amino acid metabolism
Pc21g093: weak similarity to transcription factor involved in acriflavine resistance acr-2 - Neurospora cra	0	0	0	31.0	22.9	36.8	77.8	2e-82	04 TRANSCRIPTION
Pc21g094: similarity to hypothetical transcription regulator protein - Schizosaccharomyces pombe	4e-15	SPAPB1A	gene: "SPAPB1A11.04c"; S.pombe chromosome I BAC pB1A11.	20.5	23.6	38.7	77.8	1e-106	03.01.99 other DNA processing
Pc21g174: strong similarity to zinc-finger transcription factor andA - Aspergillus nidulans	0.0	S61908	DNA-binding protein andA - Emmericella nidulans	45.6	24.9	59.1	83.8	0	0
Pc21g177: strong similarity to hellicase-like transcription factor - Homo sapiens	1e-107	ATAB5245	product: "hellicase-like transcription factor-like protein"; Arabidopsis t	32.4	16.0	28.4	28.7	1e-36	01.02.04 regulation of nitrogen and sulphur utilization
Pc21g195: similarity to cutinase transcription factor beta CTF1b - Fusarium solani	5e-24	BX897674	gene: "B2N18.280"; product: "conserved hypothetical protein"; Neur	24.1	40.0	41.8	86.5	1e-73	01.04 phosphate metabolism
Pc22g004: similarity to transcription activator Upc2 - Saccharomyces cerevisiae	4e-15	NCB9B15	gene: "B9B15.005"; product: "putative protein"; Neurospora crassa D	38.3	40.5	84.6	87.3	7e-85	01.01.04 regulation of amino acid metabolism
Pc22g018: strong similarity to SR protein kinase SkY1 - Saccharomyces cerevisiae	2e-26	T04655	protein kinase homolog FB020.10 - Arabidopsis thaliana	12.0	12.0	24.9	33.1	0.0	99 UNCLASSIFIED PROTEINS
Pc22g021: weak similarity to positive regulator of purine utilisation uuy - Aspergillus nidulans	4e-12	S54779	purine utilization positive regulator - Emmericella nidulans	12.0	12.0	17.0	42.8	1e-142	04.03.99 other tRNA-transcription activities
Pc22g125: weak similarity to transcription activator Pu3 - Saccharomyces cerevisiae	6e-23	NCAC1399	gene: "SPAC1399.05c"; S.pombe chromosome III cosmid c1399.	108.3	108.3	245.8	458.8	2e-63	01.01.04 regulation of amino acid metabolism
Pc22g160: strong similarity to Sen1 like protein An12g10480 - Aspergillus niger [putative sequencing	en5e-81	AY325411	product: "Abi-133"; Rattus norvegicus Abi-133 mRNA, complete c	21.1	18.5	43.7	48.3	1e-51	01.06.99 other lipid, fatty-acid and isoprenoid metabolism activities
Pc22g162: similarity to hypothetical transcription factor Aro80 - Saccharomyces cerevisiae	9e-85	BX649607	gene: "nira"; product: "nitrogen assimilation transcription regulator, pi	12.0	12.0	20.6	36.8	1e-135	01.05.01.01.09 aerobic aromatic catabolism
Pc24g005: weak similarity to transcription activator Upc2 - Saccharomyces cerevisiae	8e-06	S61580	probable membrane protein YDR213w - yeast (Saccharomyces cerev	15.2	12.0	21.5	28.7	1e-162	01.20.50 catabolism of secondary metabolites
Pc22g252: strong similarity to protein phosphatase methyltransferase Ppe1 - Saccharomyces cerevisiae	1e-116	BX842629	gene: "B2U13.050"; product: "related to ribosomal protein Ym52, mi	20.1	16.5	33.9	37.2	1e-168	03 CELL CYCLE AND DNA PROCESSING
Pc06g000: strong similarity to hypothetical membrane dephosphatase - Schizosaccharomyces pombe	1e-141	BX842620	gene: "B11E5.370"; product: "related to membrane dephosphatase"; Ne	88.5	78.7	157.9	174.1	1e-122	05.04 translation
Pc16g116: strong similarity to transcription factor nG1-3 like protein An11g02950 - Aspergillus niger	1e-152	NC1234A	gene: "1234A.234"; product: "related to protein 1234A.234"; Schizosac	45.1	12.0	140.8	128.1	1e-49	01.05.01 C-compound and carbohydrate utilization
Pc12g105: strong similarity to multicatalytic endopeptidase regulator like protein An11g04890 - Aspergill	1e-134	BX842627	gene: "BKJ22.150"; product: "related to GTP-binding protein 2"; Neu	138.8	127.0	221.4	268.0	0.0	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc12g110: similarity to alpha-1,6-mannosyltransferase Hoci1 - Saccharomyces cerevisiae	2e-49	AF374353	product: "putative mannosyltransferase"; Paracoccidioides brasiliensi	12.0	12.0	14.3	48.1	1e-136	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc12g146: strong similarity to hypothetical lysosomal peptidatin insensitive protease CLN2 - Canis lupus	0.0	AFU57847	gene: "tppt1"; product: "tripeptidylpeptidase 1"; Aspergillus fumigatus	23.4	16.4	51.1	84.		

Pc22g080C strong similarity to actin interacting protein like protein An02g14620 - Aspergillus niger	1e-175	BT002525 product: "Unknown protein"; Arabidopsis thaliana Unknown protein n	206.0	234.9	421.0	472.8	4e-61	06.07.99 other protein modifications
Pc22g085C weak similarity to cdc25C associated protein kinase c-tak1 - Homo sapiens	2e-09	AP006568 Gloeobacter violaceus PCC 7421 DNA, complete genome, section 1	133.2	186.7	302.7	397.1	8e-46	20 SYSTEMIC REGULATION OF / INTERACTION WITH ENVIRONMENT
Pc22g086C weak similarity to pyroglutaminyl-peptidase I PGR1 - Pyrococcus furiosus	2e-14	AK003373 Mus musculus 18-day embryo whole body cDNA, RIKEN full-length	38.4	50.0	88.9	108.0	3e-34	02.11 electron transport and membrane-associated energy conservation
Pc22g116C similarity to glutathione S-transferase omega - Sus scrofa	9e-22	ZMA01049 gene: "gstb"; product: "GSTb protein"; Zea mays mRNA for glutath	17.0	17.0	64.2	63.8	5e-48	01.01 amino acid metabolism
Pc22g145C strong similarity to protein disulfide isomerase A pdaA - Aspergillus niger	8e-34	5S7942 protein disulfide-isomerase (EC 5.3.4.1) - Aspergillus niger	407.2	182.8	823.1	169.0	3e-93	06.07 protein modification
Pc22g171C strong similarity to amidohydrolase like protein An02g0090 - Aspergillus niger	0.0	BX649605 product: "peptidase, putative"; Aspergillus fumigatus BAC pilot proje	32.0	43.7	107.8	144.4	2e-87	06.04 protein targeting, sorting and translocation
Pc22g190C strong similarity to metalloprotease like protein An04g01980 - Aspergillus niger	0.0	BX842618 gene: "B13M13.190"; product: "conserved hypothetical protein"; Neu	66.1	78.1	134.3	147.7	1e-59	01.05.04 regulation of C-compound and carbohydrate utilization
Pc24g029C strong similarity to erythrocyte splice form 1 of ankyrin like protein An12g05680 - Aspergillus	5e-33	UY084207 gene: "CG100111"; product: "SD03956P"; Drosophila melanogaster S	12.8	16.5	27.0	38.8	1e-103	01.01.07 amino acid transport
Pc12g002A strong similarity to aminotriazole resistance protein Atn1 - Saccharomyces cerevisiae	1e-55	BX649605 product: "transporter, putative"; Aspergillus fumigatus BAC pilot proje	16.8	12.0	21.3	27.8	0.0	10.05 transmembrane signal transduction
Pc12g083C strong similarity to histone shock protein Phz2 - Saccharomyces cerevisiae	1e-115	AY187281 product: "dltH peptide transporter 2"; Phaeosphaeria nodorum dltH pr	25.9	15.7	24.9	30.4	2e-48	01.05.04 regulation of C-compound and carbohydrate utilization
Pc12g123C strong similarity to anion transporter YNL275w - Saccharomyces cerevisiae	1e-130	BX842620 gene: "B11E5.170"; product: "related to chloride-bicarbonate anion ei	12.0	12.0	51.6	43.4	9e-53	01.06.07 C-compound, carbohydrate transport
Pc13g042C strong similarity to ferrioxamine B permease Srt1 - Saccharomyces cerevisiae	1e-135	S50524 hypothetical protein YEL065w - yeast (Saccharomyces cerevisiae)	58.4	52.6	77.1	115.5	1e-62	01.06.13 lipid and fatty-acid transport
Pc13g136C strong similarity to multidrug resistance protein Hdt1 - Saccharomyces cerevisiae	1e-155	T39346 probable major facilitator family multi-drug resistance protein - fission	31.9	24.8	67.2	68.7	2e-50	01.05.07 C-compound, carbohydrate transport
Pc13g138C similarity to phosphatidylinositol-phosphatidylcholine transfer protein Sec14 - Saccharomyces	8e-17	BX908789 gene: "B13D24.190"; product: "probable phosphatidylinositol/phosph	53.3	31.1	100.8	115.2	7e-43	08.16 extracellular transport, exocytosis and secretion
Pc14g0161C strong similarity to multidrug resistance protein Hdt1 - Saccharomyces cerevisiae	2e-34	S63386 HOL1 protein - yeast (Saccharomyces cerevisiae)	87.7	63.7	196.1	153.7	1e-88	99 UNCLASSIFIED PROTEINS
Pc16g034C strong similarity to hypothetical conserved protein CAD21291.1 - Neurospora crassa	1e-126	NCG51559 gene: "G15G09.040"; product: "conserved hypothetical protein"; Neur	17.7	13.7	19.9	27.8	1e-41	08.19 cellular impulse
Pc16g057C similarity to inhibitor of endosome-lysosome fusion DoxA - Legionella pneumophila	2e-24	S50468 hypothetical protein YER010c - yeast (Saccharomyces cerevisiae)	48.9	38.9	56.1	83.4	0.0	09 CELLULAR TRANSPORT AND TRANSPORT MECHANISMS
Pc16g064C similarity to high affinity copper transporter ctr4p - Schizosaccharomyces pombe	8e-27	PAN30311 gene: "ctr3"; product: "high affinity copper transporter"; Podospora a	110.3	77.0	175.1	321.0	0.0	01.06.13 lipid and fatty-acid transport
Pc16g095C strong similarity to multidrug resistance protein like protein An08g02330 - Aspergillus niger [0.0	0.0	AY246696 Alternaria brassicae ABC transporter (Atf1) gene, complete cds;	24.5	23.9	59.9	82.1	7e-95	08.13 vacuolar transport
Pc16g114C strong similarity to ATP-binding cassette multidrug transport protein atfB - Aspergillus nidula 0.0	0.0	AB060639 gene: "PMR5"; product: "ABC transporter PMR5"; Penicillium digitat	73.8	65.5	249.8	116.6	2e-52	01.05.07 C-compound, carbohydrate transport
Pc16g118C strong similarity to vacuolar H(+) Ca(2+) exchanger Vcx1 - Saccharomyces cerevisiae	2e-57	T41141 probable caCA proton/calcium exchanger - fission yeast (Schizosacc	17.4	12.8	26.9	33.8	0.0	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc16g135C similarity to multidrug resistance protein Hdt1 - Saccharomyces cerevisiae	1e-33	T39346 probable major facilitator family multi-drug resistance protein - fission	12.0	12.0	24.7	28.8	0.0	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc20g012C strong similarity to multidrug resistance protein mdr1 - Aspergillus flavus	0.0	T30882 multidrug resistance protein 1 - Aspergillus flavus	103.2	128.2	274.7	144.0	0.0	01.06.13 lipid and fatty-acid transport
Pc20g097C strong similarity to multidrug resistance protein atfB - Aspergillus nidulans [putative sequenc	0.0	AF291822 gene: "atfB"; product: "multidrug resistance protein MDR"; Trichoph	12.0	12.0	12.0	24.8	0.0	01 METABOLISM
Pc20g122C strong similarity to ATP-binding cassette multidrug transport protein atfB - Aspergillus nidula 0.0	0.0	T43022 ATP-binding multidrug cassette transport protein - Botryotinia fuckelii	12.4	30.3	51.8	25.5	2e-44	01.05.04 regulation of C-compound and carbohydrate utilization
Pc20g134C strong similarity to P-type ATPase ENA1 - Schwannomyces occidentalis	0.0	NCR24352 gene: "ena-1"; product: "putative sodium P-type ATPase"; Neurosp	12.0	12.0	18.4	33.7	0.0	14.04.03.03 pheromone response, mating-type determination, sex-specific proteins
Pc20g139C strong similarity to siderophore-iron transporter for enterobactin Ent1 - Saccharomyces cere	1e-133	AY131330 gene: "mirB"; product: "major facilitator MlRb"; Emericella nidulans n	149.7	171.6	178.2	601.1	4e-87	03.03 cell cycle
Pc20g155C strong similarity to oligopeptide transporter Opt1 - Saccharomyces cerevisiae	1e-173	T39497 hypothetical protein SPAC29b12.10c - fission yeast (Schizosacchar	24.6	26.1	38.6	56.2	2e-62	11.05.01.01 antibiotic resistance
Pc21g020C similarity to kinesin Kif21b - Mus musculus	4e-17	AF155117 product: "NY-REN-62 antigen"; Homo sapiens NY-REN-62 antigen 1	108.7	145.8	382.2	378.4	0.0	03.02 vacuolar transport
Pc21g070C strong similarity to ethionine resistance protein Ert1 - Saccharomyces cerevisiae	1e-120	AF4121H2 gene: "A1A12H2.11c"; product: "hypothetical protein"; Aspergillus fu	16.3	14.7	25.1	38.4	2e-42	01.05.07 C-compound, carbohydrate transport
Pc21g020C similarity to vacuolar H(+) Ca(2+) exchanger Vcx1 - Saccharomyces cerevisiae	2e-22	AF503229 gene: "cacx"; product: "calcium/proton exchanger"; Neurospora cras	28.5	27.0	37.0	61.1	4e-98	04 TRANSCRIPTION
Pc21g207C strong similarity to hypothetical protein B1520.50 - Neurospora crassa	1e-95	T51035 hypothetical protein B1520.50 [imported] - Neurospora crassa	87.8	122.7	176.4	236.0	1e-56	01.05.07 C-compound, carbohydrate transport
Pc21g219C strong similarity to kinesin light chain like protein An19g00300 - Aspergillus niger	5e-36	AY392413 Streptomyces clavuligerus claiin pSCL2 clone contig3.73, partial	28.9	21.2	28.7	70.2	3e-79	13.01.01.99 homeostasis of other cations
Pc22g045C similarity to fluconazole resistance transporter Flr1 - Saccharomyces cerevisiae	1e-24	T39346 probable major facilitator family multi-drug resistance protein - fission	18.7	19.4	34.3	50.4	0.0	01.05.04 regulation of C-compound and carbohydrate utilization
Pc22g046C strong similarity to iron and manganese transporter Ccc1 - Saccharomyces cerevisiae	6e-37	S43453 GC01 protein - yeast (Saccharomyces cerevisiae)	12.8	12.8	37.5	30.1	1e-31	01.05.07 C-compound, carbohydrate transport
Pc22g155C strong similarity to tetracycline resistance protein like protein An15g02680 - Aspergillus niger	1e-19	T37692 probable transport protein - fission yeast (Schizosaccharomyces pom	36.6	42.8	55.5	99.7	0.0	0.0
Pc22g241A similarity to membrane protein Tpo2 - Saccharomyces cerevisiae [truncated ORF] [putative pr	1e-18	T41018 probable membrane transporter - fission yeast (Schizosaccharomyces	12.0	16.4	27.9	27.6	0.0	0.0
Pc12g008C similarity to integral membrane protein PTH11 - Magnaporthe grisea	1e-22	AF239397 gene: "pth"; product: "integral membrane protein"; Blumeria graminis	35.9	34.9	83.4	89.3	9e-95	04.05.01.04 transcriptional control
Pc12g030C similarity to integral membrane protein PTH11 - Magnaporthe grisea	4e-23	AF119670 gene: "PTH11"; product: "integral membrane protein"; Magnaporthe	65.6	83.8	170.9	133.3	1e-60	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc12g058C similarity to vegetative incompatibility factor het-e1 - Podospora anserina [putative sequenc	1e-30	AF323583 gene: "het-e"; product: "beta transducin-like protein HET-EC2C4"; Pc	22.8	18.7	64.9	63.7	1e-69	99 UNCLASSIFIED PROTEINS
Pc13g028C strong similarity to retinol short-chain dehydrogenase like protein An19g05900 - Aspergillus	1e-79	NCB1781 gene: "B17B1.060"; product: "related to a retinol short-chain dehydro	12.0	12.5	16.3	31.7	5e-73	01.05.07 C-compound, carbohydrate transport
Pc13g047C strong similarity to nonhemolytic phospholipase like protein An01g14340 - Aspergillus niger	1e-174	AX952974 uniamed RFL: Sequence 2 from Patent WO03907825.	281.8	455.8	880.7	1279.8	5e-77	01.04 phosphate metabolism
Pc13g127C strong similarity to glucose permease Rgt2 - Saccharomyces cerevisiae	1e-156	NC13E11 gene: "13E11.140"; product: "probable sugar transporter"; Neurosp	379.8	243.9	230.5	462.8	4e-59	01.04.04 regulation of phosphate utilization
Pc16g036C weak similarity to serine protein kinase SRPK1 - Homo sapiens	2e-25	NCB1014 gene: "B1D14.120"; product: "probable dsl-1-suppressing protein kins	95.5	88.0	159.7	201.2	1e-130	10 CELLULAR COMMUNICATION/SIGNAL TRANSDUCTION MECHANISM
Pc21g000C strong similarity to histidine kinase like protein An10g00250 - Aspergillus niger	1e-110	AY456015 gene: "HHK12"; product: "putative histidine kinase HHK12p"; Cochli	12.0	12.0	22.1	29.9	1e-135	10.05 transmembrane signal transduction
Pc21g189C strong similarity to beta transducin-like protein like protein An10g1380 - Aspergillus niger	3e-20	BC043394 product: "ANKRD17 protein"; Homo sapiens ankryn repeat domain	14.0	17.0	37.4	37.1	1e-148	04.05.01.04 transcriptional control
Pc22g156C strong similarity to peptide pheromone a-factor receptor Ste3 - Saccharomyces cerevisiae	1e-35	AF308905 gene: "sls3"; product: "putative a-factor pheromone receptor Ste3";	23.0	25.4	53.1	78.8	5e-72	02.11 electron transport and membrane-associated energy conservation
Pc16g008C strong similarity to histone shock protein Hsp104 - Saccharomyces cerevisiae	0.0	AY229978 product: "Tset shock protein LPA"; Paracoccidioides brasiliensis he	211.6	279.2	464.4	371.7	7e-67	01.03 nucleotide metabolism
Pc12g111C strong similarity to antioxidant protein like protein An11g07650 - Aspergillus niger	2e-71	AF312827 gene: "TSA1"; product: "putative thiol-specific antioxidant protein Tsa	374.6	462.6	771.5	737.2	0.0	0.0
Pc13g070C weak similarity to 2-hydroxyisoflavone reductase IFRH - Arabidopsis thaliana	2e-12	AB045984 Verticillium dahliae mRNA from tomato pathotype-specific DNA fragm	31.0	29.3	51.7	77.2	0.0	0.0
Pc13g136C strong similarity to capsular associated protein CAP10 - Filobasidiella neoformans	8e-48	AF144574 gene: "CAP10"; product: "capsular associated protein"; Filobasidiell	74.5	97.4	157.5	194.6	0.0	0.0
Pc13g138C strong similarity to capsular associated protein CAP10 - Filobasidiella neoformans	1e-53	AF144574 gene: "CAP10"; product: "capsular associated protein"; Filobasidiell	64.1	25.2	85.4	81.8	1e-108	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc16g056C similarity to protease inhibitor like protein An08g01700 - Aspergillus niger	0.0	NCB1446 gene: "B1446.030"; product: "probable benzoate 4-monooxygenase c	66.1	148.2	376.8	199.0	0.0	0.0
Pc16g121C strong similarity to cytochrome P450 pisin demethylase PDA19 - Nectria haematococca	6e-48	AY220913 gene: "comD"; product: "ComD"; Dicoselotium discoidium ComD (C	122.0	98.0	331.3	409.1	2e-88	01.04.07 phosphate transport
Pc16g129C strong similarity to ABC transporter CDRA - Candida albicans	0.0	T30541 ABC1 transport protein - rice blast fungus	602.5	440.1	1101.3	1238.8	2e-88	01.01 stress response
Pc16g138C strong similarity to hypothetical membrane transport protein SPAC3H1.06c - Schizosacchar	1e-78	AY220913 gene: "comD"; product: "ComD"; Dicoselotium discoidium ComD (C	12.0	12.0	17.4	30.4	8e-54	01.05.04 regulation of C-compound and carbohydrate utilization
Pc16g142C manganese superoxide dismutase AAC35633 - Penicillium chrysogenum	1e-116	AF026523 product: "manganese superoxide dismutase"; Penicillium chrysogenu	42.5	15.2	30.0	46.3	4e-62	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc16g143C strong similarity to hypothetical membrane transport protein SPAC3H1.06c - Schizosacchar	3e-52	AF448056 gene: "dolC"; product: "putative dolichostimion transporter"; Mycospha	18.5	13.0	34.3	59.7	1e-101	01.01.01.07 biosynthesis of the aspartate family
Pc16g017C strong similarity to glutathione peroxidase Hyr1 - Saccharomyces cerevisiae	8e-63	BX842620 gene: "B11E5.410"; product: "probable glutathione peroxidase"; Neu	280.7	254.1	384.3	509.8	0.0	0.0
Pc20g068C strong similarity to D-amino-acid oxidase DAO - Fusarium solani	1e-37	AF309689 product: "putative D-amino acid oxidase G6G6.6"; Neurospora crass	29.8	24.8	62.2	79.8	5e-26	01.03.04 pyrimidine nucleotide metabolism
Pc21g027C similarity to cytochrome P450 pisin demethylase like protein An02g00150 - Aspergillus niger	0.0	0.0	12.0	12.0	29.5	39.8	5e-68	11.07 detoxification
Pc21g061C strong similarity to thioredoxin - Aspergillus nidulans	4e-48	AY49888 thioredoxin - Penicillium chrysogenum	192.2	291.2	632.9	720.1	2e-99	01 METABOLISM
Pc21g108C similarity to 7-aminoclolesterol resistance protein Rta1 - Saccharomyces cerevisiae	2e-14	NCG15D1 gene: "G15D1.020"; product: "conserved hypothetical protein"; Neur	347.4	175.0	872.2	572.7	7e-88	13.01.01.03 homeostasis of protons
Pc22g133C strong similarity to cytochrome P450 52A4 - Candida maltosa	7e-85	JS0203 cytochrome P450 52A1, alkane-inducible - yeast (Candida tropicalis)	66.3	48.0	95.1	137.9	2e-51	01.01 ionic homeostasis
Pc06g008C similarity to Na+ H+ antiporter NapA - Enterococcus hirae	4e-37	AF441858 gene: "SSC1"; product: "sodium-hydrogen antiporter"; Filobasidiell	18.1	12.0	28.7	37.1	2e-88	01.02.01.07 nitric oxide biosynthesis
Pc13g060C strong similarity to Na+ H+ exchanger protein like protein An14g01710 - Aspergillus niger	2e-50	AF441858 gene: "SSC1"; product: "sodium-hydrogen antiporter"; Filobasidiell	25.1	27.7	46.3	64.4	4e-18	01.07.10 transport of vitamins, cofactors, and prosthetic groups
Pc21g087C strong similarity to serine threonine-specific protein kinase HakA - Saccharomyces cerevisiae	4e-43	T38473 probable serine/threonine-specific protein kinase (EC 2.7.1.1) - fission	20.0	18.2	62.5	35.8	1e-68	13.01.01.03 homeostasis of protons
Pc22g146C similarity to copper transport protein like protein An08g00370 - Aspergillus niger	2e-07	AF166608 gene: "tahA"; product: "copper chaperone tahA"; Trametes versicok	23.6	48.4	93.8	74.4	0.0	0.0
Pc22g225C strong similarity to K+ H+ exchanger Kht1 - Saccharomyces cerevisiae	1e-129	T50062 Probable K+/H+-dependent [imported] - fission yeast (Schizosacchar	143.3	185.6	384.3	421.5	1e-137	02.11.05 accessory proteins of electron transport and membrane-associated energy conservation
Pc21g145C weak similarity to blastomeres yeast phase-specific protein 1 b1st - Ajellomyces dermatitidis	8e-27	NCB24G2Z gene: "B24G20.140"; product: "related to blastomeres yeast phase-s	520.7	340.6	944.1	1098.8	5e-75	02 ENERGY
Pc22g201C Pc22g20190 peroxin-1 PEX1 - Penicillium chrysogenum	0.0	AF233276 gene: "PEX1"; product: "peroxin-1"; Penicillium chrysogenum peroxi	227.5	132.6	198.8	261.6	0.0	0.0
Pc16g001C strong similarity to PRG3 protein like protein An15g07360 - Aspergillus niger	2e-92	ENCS071TX DNA centromere region sequence from BAC DP15B03, DP38F06 d	12.0	12.0	14.9	45.6	0.0	0.0
Pc22g072C strong similarity to hypothetical neutral amino acid permease - Neurospora crassa	1e-119	S47893 neutral amino acid permease - Neurospora crassa	17.4	13.8	21.2	17.4	0.0	0.0
Pc06g000C strong similarity to hypothetical protein contig12.fta_800cg - Aspergillus fumigatus	0.0	0.0	101.4	39.1	118.9	125.5	0.0	0.0
Pc06g000C strong similarity to penicilliolysin - Penicillium citrinum	1e-129	S47635 penicilliolysin - Penicillium citrinum	44.8	18.5	121.0	125.7	0.0	0.0
Pc06g006C strong similarity to hypothetical protein An08g00900 - Aspergillus niger	2e-49	BX842620 gene: "B11E5.130"; product: "hypothetical protein"; Neurospora cras	53.4	62.4	83.5	137.9	1e-64	01.01.01.01.02.05 biosynthesis of glutamate
Pc06g014C strong similarity to hypothetical protein SPAC1039.06 - Schizosaccharomyces pombe	1e-47	T50066 hypothetical protein SPAC1039.06 [imported] - fission yeast (Schizos	12.0	12.0	12.0	41.3	0.0	0.0
Pc06g015C strong similarity to hypothetical protein - Chromobacterium violaceum	2e-30	AE016910 Chromobacterium violaceum ATCC 12472 section 1 of 16 of the	12.0	12.0	29.9	27.0	0.0	0.0
Pc06g018C weak similarity to hypothetical protein PA4177 - Pseudomonas aeruginosa	4e-04	D83123 hypothetical protein PA4177 [imported] - Pseudomonas aeruginosa (C	14.3	15.8	41.7	48.7	0.0	0.0
Pc06g020C weak similarity to hypothetical protein contig1492.0.fta_2370cg - Aspergillus fumigatus	0.0	0.0	12.0	12.0	23.5	27.9	0.0	0.0
Pc06g020C weak similarity to hypothetical protein contig1492.0.fta_2370cg - Aspergillus fumigatus	0.0	0.0	12.0	12.0	23.5	27.9	0.0	0.0
Pc06g020C similarity to hypothetical protein An12g02590 - Aspergillus niger	0.0	0.0	12.0	12.0	56.7	41.4	0.0	0.0
Pc06g024C hypothetical protein	0.0	0.0	240.6	188.8	342.1	463.7	0.0	0.0
Pc08g000C strong similarity to hypothetical protein mg05551.1 - Magnaporthe grisea	0.0	0.0	17.4	20.3	21.0	59.7	1e-33	03.03.01.01.11 mitosis
Pc12g009C weak similarity to hypothetical protein BXJ22.030 - Neurospora crassa	3e-09	BX48221 gene: "BXJ22.030"; product: "conserved hypothetical protein"; Neur	24.4	24.4	38.8	48.6	0.0	0.0
Pc12g010C strong similarity to hypothetical protein An15g03240 - Aspergillus niger	0.0	0.0	57.4	88.1	136.4	81.1	1e-55	99 UNCLASSIFIED PROTEINS
Pc12g011C strong similarity to hypothetical protein contig31_part.1.fta_1450cg - Aspergillus fumigatus	5e-72	T51067 hypothetical protein B2A19.10 [imported] - Neurospora crassa	145.4	92.3	183.8	187.3	1e-70	01.04 phosphate metabolism
Pc12g017C similarity to hypothetical protein babA02b05p1 - Kluyveromyces fragilis</								

Pc12g022: hypothetical protein		0	0	0	12.0	12.0	15.9	29.6	7e-72	01 METABOLISM
Pc12g026: strong similarity to hypothetical protein mg05066.1 - Magnaporthe grisea	1e-32		BX640440	product: "putative aldolase"; Bordetella bronchiseptica strain RB50, c	12.0	12.0	12.2	42.6	0	0
Pc12g028: strong similarity to hypothetical protein 1384_scaffold_6.1fa_150cg - Fusarium graminearum	7e-13		AY228175	product: "putative arylamine N-acetyltransferase"; Streptomyces mur	16.9	24.6	82.8	77.6	0	0
Pc12g036: similarity to hypothetical protein c0146_1_165_scaffold_14.1fa_30cg - Aspergillus nidulans		0	0	0	12.0	12.0	0	24.5	0	0
Pc12g038: strong similarity to hypothetical protein - Penicillium chrysogenum	5e-28		AB091505	product: "hypothetical protein"; Penicillium chrysogenum mRNA for	12.0	12.0	22.4	29.3	1e-63	01 METABOLISM
Pc12g060: strong similarity to hypothetical protein An07g02930 - Aspergillus niger		0	0	0	30.6	31.8	123.4	109.8	9e-32	01 METABOLISM
Pc12g063: strong similarity to hypothetical oxidoreductase of the short-chain dehydrogenase reductase f1e-51			E87298	hypothetical protein CC0398 [Imported] - Caulobacter crescentus	27.1	36.6	62.9	55.6	0	0
Pc12g068: strong similarity to hypothetical protein Ta0729 - Thermoplasma acidophilum	6e-22		TACID3_8	gene: "Ta0729"; product: "conserved hypothetical protein"; Thermop	314.5	325.8	726.1	684.1	0	0
Pc12g069: strong similarity to hypothetical protein An01g0800 - Aspergillus niger	7e-86		ENAC133	product: "ORF"; Emericella nidulans cosmid SW06E08, complete s	25.9	18.4	31.7	53.1	1e-103	11.01 stress response
Pc12g069: strong similarity to hypothetical protein contig1477_1.1fa_810cg - Aspergillus fumigatus		0	0	0	93.7	93.8	293.9	259.1	9e-79	03.01.05 DNA recombination and DNA repair
Pc12g078: strong similarity to hypothetical protein SPCPC757.02c - Schizosaccharomyces pombe	5e-60		T41593	hypothetical protein SPCPC757.02c - fission yeast (Schizosaccharom	12.0	12.0	44.1	56.7	0	0
Pc12g080: strong similarity to siruin type 5 SIRT5 - Homo sapiens	2e-41		AE010224	gene: "PF1154"; product: "transcriptional regulatory protein, sir2"; P	94.9	84.7	98.0	172.5	0	0
Pc12g081: strong similarity to hypothetical protein contig1492_0.1fa_2200cg - Aspergillus fumigatus	4e-09		NCB11N2	gene: "B11N2.100"; product: "putative protein"; Neurospora crassa D	22.9	19.0	63.7	44.8	3e-58	01.01.04 regulation of amino acid metabolism
Pc12g086: strong similarity to hypothetical conserved protein 3H10.120 - Neurospora crassa	9e-96		NC3H10_1	gene: "3H10.120"; product: "conserved hypothetical protein"; Neuros	24.7	34.2	83.3	93.8	0	0
Pc12g087: weak similarity to transcription regulator of maltose utilization amyR - Aspergillus oryzae	3e-07		ANU5099	gene: "lacB"; product: "FacB"; Aspergillus niger putative DNA bindin	69.5	63.7	138.1	148.2	1e-148	04.05.01.04 transcriptional control
Pc12g087: strong similarity to hypothetical protein H0753 - Synechocystis sp.	6e-66		AE012994	gene: "TTE0217"; product: "conserved hypothetical protein"; Therm	120.0	120.0	21.3	27.1	1e-58	01.01.01 amino acid biosynthesis
Pc12g087: strong similarity to conserved hypothetical protein - Xanthomonas axonopodis	1e-79		AE012042	gene: "XAC3929"; product: "conserved hypothetical protein"; Xantho	827.9	670.2	2649.0	2364.8	2e-72	01.06.01.05 fatty acid biosynthesis
Pc12g090: strong similarity to hypothetical protein contig1496_7.1fa_220cg - Aspergillus fumigatus	8e-38		AE016924	Chromobacterium violaceum ATCC 12472 section 15 of 16 of the	38.8	62.4	100.9	102.2	5e-43	01.02 nitrogen and sulfur metabolism
Pc12g0917: similarity to hypothetical protein SMb20006 - Sinorhizobium meliloti	1e-32		AE016778	product: "oxidoreductase, FAD-binding"; Pseudomonas putida KT24-	63.9	56.5	164.3	190.9	0	0
Pc12g094: similarity to conserved hypothetical protein - Bacillus anthracis	1e-12		AE017030	product: "conserved hypothetical protein"; Bacillus anthracis str. Ame	70.0	89.5	131.0	207.3	0	0
Pc12g095: similarity to hypothetical protein An17g00600 - Aspergillus niger		0	0	0	24.2	16.1	29.8	39.6	0	0
Pc12g101: strong similarity to hypothetical protein contig_1_86_scaffold_5.1fa_140cg - Aspergillus nidu		0	0	0	17.5	12.0	73.0	50.8	3e-73	01.02.01.07 nitric oxide biosynthesis
Pc12g105: strong similarity to hypothetical protein contig1496_0.1fa_1440cg - Aspergillus fumigatus		0	0	0	38.8	39.8	77.0	88.4	9e-40	01.01.04 regulation of amino acid metabolism
Pc12g107: similarity to hypothetical protein 1168_scaffold_2.1fa_590cg - Fusarium graminearum	4e-12		AY437641	gene: "ST1"; product: "serine/threonine kinase"; Leptothorax macr	36.8	21.1	46.6	47.1	0	0
Pc12g111: weak similarity to hypothetical transcription regulator SPAC1103.07c - Schizosaccharomyces	7e-06		S48917	probable regulatory protein YHR178w - yeast (Saccharomyces cervi	18.9	36.9	44.8	60.4	1e-155	99 UNCLASSIFIED PROTEINS
Pc12g113: hypothetical protein		0	0	0	58.3	60.5	310.6	143.2	9e-27	01.01.01.01.02 biosynthesis of the glutamate group (proline, hydroxyprolin, arginine, glutamine, glutamate)
Pc12g124: strong similarity to hypothetical protein contig1490_3.1fa_450cg - Aspergillus fumigatus	6e-48		AE016921	Chromobacterium violaceum ATCC 12472 section 12 of 16 of the	50.1	37.2	71.3	77.4	1e-61	99 UNCLASSIFIED PROTEINS
Pc12g124: strong similarity to hypothetical protein 1141_scaffold_2.1fa_370cg - Fusarium graminearum	4e-30		NC80A10	gene: "80A10.200"; product: "conserved hypothetical protein"; Neut	12.0	12.0	46.3	43.8	3e-71	01.05 C-compound and carbohydrate metabolism
Pc12g127: similarity to hypothetical protein YPL067c - Saccharomyces cerevisiae	7e-12		S60923	hypothetical protein YPL067c - yeast (Saccharomyces cerevisiae)	107.5	63.8	156.2	137.7	0	0
Pc12g132: strong similarity to hypothetical protein contig46_part_1.1fa_3260cg - Aspergillus fumigatus	2e-72		AE012994	gene: "TTE0212"; product: "predicted sugar phosphate isomerase";	466.4	540.0	863.9	1413.8	0	0
Pc12g145: strong similarity to hypothetical protein contig_1_55_scaffold_3.1fa_710cg - Aspergillus nidul	5e-63		AE016790	product: "polysaccharide deacetylase family protein"; Pseudomonas	12.0	12.0	14.1	35.0	0	0
Pc12g145: hypothetical protein		0	0	0	104.5	102.4	407.6	503.0	0	0
Pc12g147: similarity to hypothetical protein contig_1_122_scaffold_9.1fa_160cg - Aspergillus nidulans		0	0	0	549.8	285.9	970.7	1130.0	0.0	01.06.13 lipid and fatty-acid transport
Pc12g147: hypothetical protein		0	0	0	17.9	12.0	47.3	43.8	0	0
Pc12g158: strong similarity to hypothetical protein An12g05670 - Aspergillus niger		0.0	ST7690	probable membrane protein YOLO75c - yeast (Saccharomyces cervi	18.4	19.2	31.5	58.4	0	0
Pc12g002: strong similarity to hypothetical protein An05g00170 - Aspergillus niger		0	0	0	44.4	86.9	111.9	133.1	0	0
Pc13g005: similarity to hypothetical protein ncu01851.1 - Neurospora crassa		0	0	0	12.2	12.0	16.8	27.1	4e-93	01 METABOLISM
Pc13g006: weak similarity to hypothetical protein BAB55393.1 - Homo sapiens		0	0	0	36.2	23.3	55.0	91.3	1e-144	11.01 stress response
Pc13g015: strong similarity to hypothetical protein contig589.1fa_450cg - Aspergillus fumigatus	9e-52		S57117	probable aldehyde reductase (EC 1.1.1.1) - YJR096w - yeast (Sacchar	34.2	42.7	67.9	123.3	0	0
Pc13g028: similarity to hypothetical protein SPCPC757.02c - Schizosaccharomyces pombe	2e-37		NCB9811	gene: "B9B11.170"; product: "conserved hypothetical protein"; Neut	12.0	12.0	12.0	26.8	3e-78	01.05.01 C-compound and carbohydrate utilization
Pc13g028: hypothetical protein		0	0	0	12.0	12.0	24.1	24.1	0	0
Pc13g036: strong similarity to hypothetical protein contig31_part_1.1fa_1050cg - Aspergillus fumigatus	5e-30		AB009053	Arabidopsis thaliana genomic DNA, chromosome 5, P1 clone:MQB2	162.5	182.1	337.2	333.1	0	0
Pc13g039: strong similarity to hypothetical protein An12g00900 - Aspergillus niger		0	0	0	20.9	17.5	32.0	36.8	6e-38	10.01 intracellular signalling
Pc13g042: strong similarity to hypothetical protein An15g04720 - Aspergillus niger		0	0	0	71.8	62.0	111.1	132.0	0	0
Pc13g047: strong similarity to hypothetical protein An05g00690 - Aspergillus niger	4e-05		BC057529	gene: "supt5h"; product: "supt5h protein"; Danio rerio suppressor of	364.7	417.0	820.4	513.0	0	0
Pc13g047: strong similarity to hypothetical protein An08g01950 - Aspergillus niger	2e-87		AE015942	gene: "CTC01793"; product: "transcriptional regulator"; Clostridium	96.4	105.3	344.5	325.8	9e-57	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc13g048: strong similarity to hypothetical protein An05g00625 - Aspergillus niger	6e-04		BC041788	product: "Similar to RIKEN cDNA 111002S15 gene"; Homo sapiens	114.3	73.8	172.0	155.9	0	0
Pc13g048: strong similarity to hypothetical protein contig_1_100_scaffold_7.1fa_370cg - Aspergillus nid	7e-32		BC038340	product: "dehydrogenase/reductase (SDS family) member 8"; Mus m	43.2	18.6	66.8	112.3	1e-126	30.16 mitochondrion
Pc13g049: hypothetical protein		0	0	0	12.0	14.5	35.1	32.2	0	0
Pc13g055: strong similarity to hypothetical protein mg06134.1 - Magnaporthe grisea	3e-49		T40474	hypothetical protein SPBC484.02c - fission yeast (Schizosaccharom	71.9	74.1	121.8	172.5	0	0
Pc13g061: similarity to hypothetical intracellular protease amidase related enzyme of the ThiJ family CA(2e-16			G97247	intracellular protease/amidase related enzyme (ThiJ family) [Imported	34.9	23.4	85.3	71.0	0	0
Pc13g063: strong similarity to hypothetical protein contig1477_1.1fa_530cg - Aspergillus fumigatus	7e-06		T38995	hypothetical protein SPAC635.03c - fission yeast (Schizosaccharomy	332.5	302.0	451.1	633.8	2e-89	01.02.01 nitrogen and sulfur utilization
Pc13g072: strong similarity to hypothetical protein contig1495_2.1fa_590cg - Aspergillus fumigatus	8e-19		BX640418	product: "conserved hypothetical protein"; Bordetella pertussis strain	31.2	52.6	66.9	120.7	0	0
Pc13g074: strong similarity to hypothetical protein contig1488_2.1fa_790cg - Aspergillus fumigatus	1e-104		BX424918	gene: "B13M13.090"; product: "conserved hypothetical protein"; Neu	34.3	61.9	76.1	135.3	1e-52	99 UNCLASSIFIED PROTEINS
Pc13g074: strong similarity to hypothetical protein An12g02830 - Aspergillus niger		0	0	0	12.0	13.9	32.6	51.1	1e-151	40 SUBCELLULAR LOCALISATION
Pc13g078: similarity to hypothetical protein YER080w - Saccharomyces cerevisiae	2e-13		S05083	hypothetical protein YER080w - yeast (Saccharomyces cerevisiae)	26.6	16.5	30.3	37.1	2e-40	99 UNCLASSIFIED PROTEINS
Pc13g089: strong similarity to hypothetical protein SPBC354.08c - Schizosaccharomyces pombe	1e-116		T40288	hypothetical protein SPBC354.08c - fission yeast (Schizosaccharom	137.0	114.1	227.9	246.0	0	0
Pc13g099: weak similarity to hypothetical protein YGL079w - Saccharomyces cerevisiae	8e-22		NCB9811	gene: "B9B11.220"; product: "conserved hypothetical protein"; Neut	26.4	37.8	97.2	79.1	0	0
Pc13g102: hypothetical protein		0	0	0	55.6	85.8	117.1	98.9	0	0
Pc13g109: hypothetical protein		0	0	0	27.1	28.3	63.4	72.8	0	0
Pc13g111: weak similarity to hypothetical protein mg10737.1 - Magnaporthe grisea		0	0	0	12.0	17.2	18.5	36.8	6e-35	99 UNCLASSIFIED PROTEINS
Pc13g112: strong similarity to hypothetical protein An02g02460 - Aspergillus niger	3e-04		AY228175	gene: "kinU"; product: "LanU-like protein"; Streptomyces murayamae	40.3	68.1	122.3	159.2	1e-76	01.02.04 regulation of nitrogen and sulphur utilization
Pc13g112: weak similarity to hypothetical protein YER182w - Saccharomyces cerevisiae	2e-06		AX119091	unnamed ORF; Sequence 255 from Patent WO0129221.	12.0	17.2	26.9	49.5	2e-62	01.05.04 regulation of C-compound and carbohydrate utilization
Pc13g118: strong similarity to DCG1-like protein, putative - Aspergillus fumigatus	4e-80		BX649606	product: "DCG1-like protein, putative"; Aspergillus fumigatus BAC pil	19.1	23.1	30.9	46.9	7e-96	01.01 amino acid metabolism
Pc13g122: strong similarity to hypothetical protein SPAC111H1.03c - Schizosaccharomyces	5e-84		BX649606	product: "smr family protein, putative"; Aspergillus fumigatus BAC pil	180.1	169.4	254.6	346.7	0	0
Pc13g131: strong similarity to hypothetical conserved gl_19075788 - Schizosaccharomyces pombe	1e-115		T41346	probable glutamate-L-semialdehyde aminotransferase - fission yeast	57.5	37.2	106.8	239.1	1e-127	99 UNCLASSIFIED PROTEINS
Pc13g133: hypothetical protein		0	0	0	50.0	44.7	161.3	85.9	3e-37	01.03 nucleotide metabolism
Pc13g134: strong similarity to hypothetical protein SPAC1039.06c - Schizosaccharomyces pombe	7e-51		T50056	hypothetical protein SPAC1039.06 [Imported] - fission yeast (Schizot	46.9	53.3	97.5	114.4	0	0
Pc13g138: strong similarity to hypothetical protein 1348_scaffold_5.1fa_420cg - Fusarium graminearum	6e-56		T49704	hypothetical protein B23L21.70 [Imported] - Neurospora crassa	83.6	52.2	127.2	180.1	0	0
Pc13g141: strong similarity to hypothetical protein An15g05550 - Aspergillus niger		0	0	0	18.4	18.8	29.3	46.9	2e-95	11.05.01 resistance proteins
Pc13g141: strong similarity to hypothetical protein contig61_1fa_890cg - Aspergillus fumigatus		0	0	0	77.6	93.7	280.1	199.9	0	0
Pc13g143: similarity to hypothetical beta-lactamase XF1621 - Xylella fastidiosa	2e-74		NC104H10	gene: "104H10.010"; product: "conserved hypothetical protein"; Neu	77.1	49.1	120.9	116.8	2e-43	99 UNCLASSIFIED PROTEINS
Pc13g147: weak similarity to hypothetical protein B208.260 - Neurospora crassa		0	0	0	25.2	25.1	91.7	88.1	0	0
Pc13g149: strong similarity to hypothetical protein ncu04039.1 - Neurospora crassa	2e-22		BX897678	gene: "B10H18.160"; product: "conserved hypothetical protein"; Neu	49.1	50.3	164.6	225.2	8e-53	01.03 nucleotide metabolism
Pc13g152: strong similarity to hypothetical protein An15g02470 - Aspergillus niger		0	0	0	51.4	90.5	122.1	126.8	0	0
Pc14g002: strong similarity to ketoreductase krd - Zygosaccharomyces rouxi	1e-52		T38902	probable cinnamoyl-coa reductase - fission yeast (Schizosaccharom	12.0	12.0	12.0	24.9	1e-102	01 METABOLISM
Pc14g007: similarity to hypothetical protein encoded by SMC00431 - Sinorhizobium meliloti	1e-13		AP005085	gene: "VPA0515"; product: "hypothetical protein"; Vibrio parahaemol	12.0	12.0	28.4	22.1	1e-62	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc14g012: strong similarity to hypothetical 2-hydroxyhepta-2,4-diene-1,7-dioate isomerase BH2000 - BaSe55			BC009403	product: "Unknown (protein for MGC:15165)"; Homo sapiens, clone I	269.1	364.3	481.4	807.8	1e-78	01.01.01.15 biosynthesis of the pyruvate family (alanine, isoleucine, leucine, valine) and D-alanine
Pc14g018: strong similarity to hypothetical protein contig_1_98_scaffold_6.1fa_1140cg - Aspergillus nid	3e-18		U94886_1	Brachyspira hyodysenteriae ACP reductase (fabG); acyl carrier	167.0	157.2	220.3	513.8	6e-33	47.03.01.01.99 other brain areas
Pc15g001: strong similarity to hypothetical protein An02g03640 - Aspergillus niger	1e-17		AE014397	gene: "BR0950"; product: "oxidoreductase, putative"; Brucella suis 1	100.3	124.8	238.2	330.5	1e-51	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc15g006: weak similarity to hypothetical protein SPBC31F10.02 - Schizosaccharomyces pombe	8e-10		CEC25H3	gene: "C25H3.3"; product: "Hypothetical protein C25H3.3"; Caenor	145.4	121.4	172.3	374.7	3e-62	01 METABOLISM
Pc15g009: similarity to hypothetical short chain alcohol dehydrogenase cta - Cucumis sativus	3e-17		AE003507	gene: "scu"; product: "CG7113-PA"; Drosophila melanogaster chron	39.8	12.0	34.1	75.7	0	0
Pc15g010: strong similarity to hypothetical protein F49E12.10 - Caenorhabditis elegans	6e-36		T22443	hypothetical protein F49E12.9 - Caenorhabditis elegans	43.7	15.2	18.4	118.4	0	0
Pc15g013: weak similarity to hypothetical protein An05g00600 - Aspergillus niger		0	0	0	14.9	18.8	44.0	33.7	8e-81	01.01.99 other amino acid metabolism activities
Pc15g014: strong similarity to hypothetical protein An08g0840 - Aspergillus niger		0	0	0	66.0	87.2	114.4	6e-82	0	0
Pc15g018: strong similarity to hypothetical oxidoreductase PA0147 - Pseudomonas aeruginosa	3e-63		C83628	probable oxidoreductase PA0147 [Imported] - Pseudomonas aerugin	12.0	20.0	26.0	47.3	2e-55	01.03 nucleotide metabolism
Pc15g021: strong similarity to hypothetical protein encoded by CG16717 - Drosophila melanogaster	4e-32		AE003551	product: "CG16717-PA"; Drosophila melanogaster chromosome 3L	16.3	12.0	33.3	28.7	2e-12	99 UNCLASSIFIED PROTEINS
Pc16g001: strong similarity to ketoreductase krd - Zygosaccharomyces rouxi	8e-55		T38902	probable cinnamoyl-coa reductase - fission yeast (Schizosaccharom	32.0	44.7	112.4	90.4	0	0
Pc16g002: weak similarity to hypothetical protein YGR273c - Saccharomyces cerevisiae	8e-04		T39810	hypothetical protein SPBC19C7.04c - fission yeast (Schizosaccharo	66.9	32.2	85.9			

Pc16g0071 strong similarity to hypothetical protein contig50.1fa_130wg - Aspergillus fumigatus	1e-32	BC054138 product: "Unknown (protein for MGC:63829)"; Danio rerio cDNA clone	69.2	19.1	78.4	98.0	0	
Pc16g0093 strong similarity to hypothetical protein contig10.1fa_380cg - Aspergillus fumigatus	0	0	68.6	13.4	129.5	92.2	0	
Pc16g0303 strong similarity to hypothetical protein contig1492_0.1fa_6350cg - Aspergillus fumigatus	1e-179	AB110615 gene: "sdn1"; product: "saponin hydrolase precursor"; Neocostomopsis	16.1	12.0	71.3	49.6	0	
Pc16g0371 strong similarity to hypothetical protein An1700600 - Aspergillus niger	0	0	47.2	23.4	112.4	72.1	0	
Pc16g0444 strong similarity to hypothetical protein mg01514.1 - Magnaporthe grisea	0	0	41.4	93.8	155.6	79.8	0	
Pc16g0497 strong similarity to hypothetical protein 1A9.40 - Neurospora crassa	1e-107	T48700 hypothetical protein 1A9.40 [imported] - Neurospora crassa	28.2	15.8	83.1	74.4	0	
Pc16g0505 strong similarity to hypothetical protein contig46_part.1fa_950wg - Aspergillus fumigatus	7e-70	AE015942 gene: "CTC01794"; product: "transcriptional regulator"; Clostridium t	49.8	66.1	130.6	160.2	0	
Pc16g0573 strong similarity to hypothetical protein encoded by Smb20234 - Sinorhizobium meliloti	6e-58	A95870 hypothetical protein [imported] - Sinorhizobium meliloti (strain 1021) n	15.5	12.0	24.6	26.1	0	
Pc16g0611 similarity to hypothetical protein contig46_part.1fa_330cg - Aspergillus fumigatus	0	0	16.2	12.0	29.7	38.1	3e-82	01.05.07 C-compound, carbohydrate transport
Pc16g0616 weak similarity to hypothetical serine-rich protein SPCC1322.10 - Schizosaccharomyces pombe	0	0	163.9	207.9	334.0	318.1	0	
Pc16g0741 strong similarity to hypothetical protein 1323_scaffold.5.1fa_330wg - Fusarium graminearum	1e-103	T51198 hypothetical serine-rich protein - fission yeast (Schizosaccharomyces related to myo-inositol transport protein ITR1 [imported] - Neurospora	255.4	184.8	1059.9	670.9	0	
Pc16g0765 strong similarity to hypothetical protein An04g03340 - Aspergillus niger	0	0	83.8	54.2	180.4	155.6	0.0	99 UNCLASSIFIED PROTEINS
Pc16g0803 strong similarity to hypothetical protein SCF91.02c - Streptomyces coelicolor	1e-175	SCO9391 gene: "SCO0642"; "SCF91.02c"; product: "conserved hypothetical pr	81.6	108.2	189.1	243.9	8e-58	01.05 C-compound and carbohydrate metabolism
Pc16g0805 strong similarity to hypothetical protein - Streptomyces avermitilis	0.0	AP005048 product: "hypothetical protein"; Streptomyces avermitilis genomic DN	19.9	24.0	33.2	84.2	1e-99	01.05 C-compound and carbohydrate metabolism
Pc16g0823 strong similarity to hypothetical myoinositol-dehydrogenase spcB - Streptomyces spectabilis	1e-35	AF107704 gene: "spcB"; product: "SpcB"; Streptomyces spectabilis SpcC (spc	71.4	68.2	173.9	218.0	0	
Pc16g0914 strong similarity to hypothetical protein PA5145 - Pseudomonas aeruginosa	1e-123	BX087675 gene: "B2E7.160"; product: "conserved hypothetical protein"; Neuros	100.6	152.5	194.3	381.7	0	
Pc16g0955 strong similarity to hypothetical protein 1244_scaffold.3.1fa_50cg - Fusarium graminearum	1e-06	AF451898 gene: "orf128"; product: "tetracycline-resistance protein"; Helicobac	27.6	17.7	50.4	92.8	0	
Pc16g0965 strong similarity to hypothetical protein contig50.1fa_290cg - Aspergillus fumigatus	0	0	64.0	64.4	90.6	135.3	0.0	99 UNCLASSIFIED PROTEINS
Pc16g1003 hypothetical protein	0	0	27.2	32.5	75.6	102.6	0	
Pc16g1065 strong similarity to hypothetical protein YIR007w - Saccharomyces cerevisiae	1e-150	S50878 hypothetical protein YIR007w - yeast (Saccharomyces cerevisiae)	93.4	27.2	165.5	198.2	0	
Pc16g1133 strong similarity to hypothetical protein contig1490_3.1fa_1270wg - Aspergillus fumigatus	0	0	63.9	34.2	169.1	197.6	0	
Pc16g1136 strong similarity to hypothetical protein contig_1_116_scaffold.8.1fa_60wg - Aspergillus nidul	0	0	12.0	12.0	36.9	38.1	0	
Pc16g1141 strong similarity to hypothetical protein nc060371.1 - Neurospora crassa [putative sequenc	3e-07	T52364 hypothetical protein B11E6.130 [imported] - Neurospora crassa	12.0	12.0	44.3	38.1	0	
Pc16g1165 strong similarity to hypothetical protein contig46_part.1fa_1620wg - Aspergillus fumigatus	0	0	25.3	12.2	36.9	53.4	0	
Pc16g1223 similarity to hypothetical protein AAM35689.1 - Xanthomonas axonopodis	1e-15	AP006577 Gloeobacter violaceus PCC 7421 DNA, complete genome, section 1f	27.2	12.0	18.7	24.6	0	
Pc16g1243 weak similarity to hypothetical protein An01g07060 - Aspergillus niger	0	0	12.0	12.0	23.1	49.8	0	
Pc16g1303 strong similarity to hypothetical protein contig1492_0.1fa_6240cg - Aspergillus fumigatus	2e-48	AP003004 gene: "m14579"; product: "2-haloalkanoic acid dehalogenase"; Meso	27.4	42.9	70.1	74.8	1e-72	01.04 phosphate metabolism
Pc16g1323 strong similarity to hypothetical protein contig46_part.1fa_1200wg - Aspergillus fumigatus	0	0	12.0	12.0	27.2	42.9	0	
Pc16g1343 weak similarity to hypothetical protein An01g12910 - Aspergillus niger	4e-06	AF421215 gene: "snf"; product: "haloalkanoic acid dehalogenase"; Colletotrichum gloeospor	45.7	47.8	122.4	116.8	9e-40	01.05.01.01 C-compound, carbohydrate catabolism
Pc16g1373 similarity to hypothetical protein contig46_part.1fa_1100wg - Aspergillus fumigatus	0	0	259.1	162.0	462.8	372.9	0	
Pc16g1423 strong similarity to hypothetical protein contig5_part.1fa_940wg - Aspergillus fumigatus	5e-16	AF409109 product: "14.5 kDa bacteriolytic enzyme"; Dermatoaphidius pterony	12.0	12.0	53.9	58.9	0	
Pc16g1443 similarity to hypothetical protein contig1477_1.1fa_1040wg - Aspergillus fumigatus	0	0	50.1	77.1	176.0	159.1	0	
Pc16g1465 strong similarity to hypothetical protein contig_1_139_scaffold.11.1fa_640wg - Aspergillus ni	0	0	821.8	442.1	1648.8	2938.4	0	
Pc16g1483 similarity to hypothetical protein contig46_part.1fa_2020cg - Aspergillus fumigatus	0	0	24.5	42.2	83.8	37.2	0	
Pc16g1525 similarity to hypothetical protein An08g01250 - Aspergillus niger	0	0	149.8	150.5	365.8	385.8	0	
Pc16g1533 similarity to hypothetical protein 1367_scaffold.6.1fa_40cg - Fusarium graminearum	0	0	325.9	311.3	556.0	678.9	0	
Pc16g1533 strong similarity to hypothetical protein An12g10330 - Aspergillus niger	0	0	16.4	19.4	40.7	45.9	0	
Pc16g1533 similarity to hypothetical protein nc05318.1 - Neurospora crassa	0	0	201.6	204.5	414.7	532.7	0	
Pc16g1533 weak similarity to hypothetical protein An03g00300 - Aspergillus niger	0	0	22.3	16.3	49.0	41.8	0	
Pc16g1544 strong similarity to hypothetical protein An01g10730 - Aspergillus niger	8e-44	AC115592 product: "hypothetical protein"; Dictyostelium discoideum chromosom	364.2	655.1	958.4	966.1	0	
Pc17g0011 hypothetical protein	0	0	152.0	118.1	476.4	427.8	0	
Pc18g0013 similarity to integral membrane protein PTH11 - Magnaporthe grisea [putative sequencing err	1e-16	NC94C8_2 gene: "94C8.020"; product: "related to integral membrane protein PTI	43.9	106.2	152.6	88.9	1e-118	01.01.01 amino acid biosynthesis
Pc18g0014 strong similarity to hypothetical conserved protein 99H12.80 - Neurospora crassa	3e-93	BX649605 product: "hypothetical protein, conserved"; Aspergillus fumigatus BA	72.4	141.3	228.4	309.4	0	
Pc18g0031 strong similarity to hypothetical protein CP0630 - Chlamydomonas reinhardtii	1e-114	NCB1383.3 gene: "B1383.110"; product: "conserved hypothetical protein"; Neur	12.0	12.0	12.0	34.1	0	
Pc18g0033 strong similarity to hypothetical protein contig336.1fa_290wg - Aspergillus fumigatus [putative	0	0	12.0	12.0	27.5	26.1	1e-51	01.03 nucleotide metabolism
Pc18g0051 similarity to hypothetical phosphoglycerate mutase SPAC5H10.03 - Schizosaccharomyces p	2e-20	NC49D12_2 gene: "49D12.140"; product: "conserved hypothetical protein"; Neur	12.0	12.0	14.5	45.2	1e-110	40.30 prokaryotic cell membrane (inner membrane of gram - bacteria)
Pc18g0105 strong similarity to hypothetical UDP-glucose 4-epimerase - Alcaligenes eutrophus	4e-63	BX649607 product: "possible epimerase"; Aspergillus fumigatus BAC pilot proj	16.6	22.2	27.0	59.1	0.0	01.05.01.01 C-compound, carbohydrate catabolism
Pc18g0113 similarity to hypothetical protein Aul4 - Saccharomyces cerevisiae	2e-19	S19386 hypothetical protein YOL1038c - yeast (Saccharomyces cerevisiae)	166.6	241.9	360.0	566.1	2e-50	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc18g0131 strong similarity to hypothetical beta-hexosaminidase A precursor BH0675 - Bacillus halodurans	4e-91	AP004604 gene: "OB3206"; product: "beta-N-acetylhexosaminidase (beta-hexos	131.4	173.2	283.8	363.7	0	
Pc18g0133 strong similarity to hypothetical protein contig_1_172_scaffold.16.1fa_550cg - Aspergillus n	2e-55	AF061253 gene: "GST1"; product: "glutathione S-transferase"; Botryotinia fucki	563.9	677.7	1417.0	1418.4	0	
Pc18g0184 strong similarity to hypothetical methyltransferase AAO34671.1 - Gibberella zeae	4e-75	AF359360 gene: "OrfF"; product: "putative methyltransferase"; Fusarium sporot	802.0	768.7	2089.2	2059.1	9e-45	03.03.01.01.11 mitosis
Pc18g0245 strong similarity to hypothetical protein contig_1_66_scaffold.4.1fa_40cg - Aspergillus nidul	0	0	12.0	13.3	32.8	47.8	5e-70	01 METABOLISM
Pc18g0265 strong similarity to hypothetical protein contig5_part.1fa_90cg - Aspergillus fumigatus	6e-30	AB107895 Aspergillus kawachii cwpB gene for hypothetical protein, complete	1790.1	2021.9	4263.9	3792.8	4e-73	04.05.01.04 transcriptional control
Pc18g0271 strong similarity to hypothetical protein contig1488_2.1fa_1170cg - Aspergillus fumigatus	8e-81	AF102927 gene: "T3A1"; product: "putative thiol-specific antioxidant protein Tsa1	1291.4	1848.8	3341.5	3891.1	1e-167	08 CELLULAR TRANSPORT AND TRANSPORT MECHANISMS
Pc18g0284 strong similarity to hypothetical protein contig_1_10_scaffold.1.1fa_200cg - Aspergillus nidul	5e-24	AC002994 gene: "m10013"; product: "transcriptional regulator"; Mesorhizobium	15.7	23.3	37.9	51.8	1e-38	05.01 ribosome biogenesis
Pc18g0303 strong similarity to hypothetical protein contig_1_167_scaffold.14.1fa_30wg - Aspergillus nid	2e-44	NTG1_MU SODIUM- AND CHLORIDE-DEPENDENT GABA TRANSPORTER	89.6	58.3	107.2	224.1	9e-49	99 UNCLASSIFIED PROTEINS
Pc18g0533 strong similarity to hypothetical protein contig248.1fa_180wg - Aspergillus fumigatus	4e-31	CAL39049 gene: "rps21"; product: "ribosomal protein S21"; Candida albicans m	59.4	50.8	86.2	120.1	1e-77	01.03.01.01 purine nucleotide catabolism
Pc18g0594 weak similarity to hypothetical protein T17H3.1 - Arabidopsis thaliana	1e-05	AY045812 gene: "At1g27510"; product: "unknown protein"; Arabidopsis thaliana	12.8	12.0	27.7	34.6	1e-116	08.16 extracellular transport, exocytosis and secretion
Pc20g0055 strong similarity to hypothetical protein nc01441.1 - Neurospora crassa	1e-64	NC820D11 gene: "B20D17.150"; product: "probable inosine triphosphate pyropho	200.8	168.7	291.2	342.6	0	
Pc20g0085 strong similarity to hypothetical protein SPBC27812.12c - Schizosaccharomyces pombe	1e-115	SPAB4538 gene: "pil66"; product: "probable membrane protein YOL130w"; Sci	73.3	74.3	150.1	103.2	0	
Pc20g0143 strong similarity to hypothetical protein An12g06840 - Aspergillus niger	0	0	101.4	81.0	148.5	171.2	0	
Pc20g0185 strong similarity to hypothetical protein SC9B2.03 - Streptomyces coelicolor	1e-169	T35899 hypothetical protein SC9B2.03 - Streptomyces coelicolor	16.0	16.5	52.4	76.6	0	
Pc20g0203 strong similarity to hypothetical protein mg01347.1 - Magnaporthe grisea	7e-43	NC80A10_2 gene: "80A10.320"; product: "conserved hypothetical protein"; Neur	26.1	28.2	70.6	60.9	8e-91	01.01.04 regulation of amino acid metabolism
Pc20g0211 strong similarity to hypothetical protein An02g13400 - Aspergillus niger	0	0	12.0	12.0	32.2	34.3	5e-23	04 TRANSCRIPTION
Pc20g0235 weak similarity to hypothetical transcription regulator SPBC530.05 - Schizosaccharomyces p	6e-10	T49482 hypothetical protein B14D6.340 [imported] - Neurospora crassa	15.0	12.0	15.0	24.6	0	
Pc20g0351 strong similarity to hypothetical protein contig_1_51_scaffold.3.1fa_1450cg - Aspergillus nid	2e-14	AX885834 unnamed ORF; Sequence 1697 from Patent EP1033401.	127.6	165.3	589.7	344.8	0	
Pc20g0363 similarity to integral membrane protein PTH11 - Magnaporthe grisea	5e-20	NCB7H23_3 gene: "B7H23.130"; product: "related to L-fucose permease"; Neuro	12.0	12.0	38.1	36.8	9e-43	06.13 proteolytic degradation
Pc20g0391 weak similarity to hypothetical protein YBR246w - Saccharomyces cerevisiae	8e-39	CNS09545 DNA centromeric region sequence from BAC DP26B06, DP34F04, I	19.2	20.6	115.1	54.7	1e-56	01.04 phosphate metabolism
Pc20g0405 similarity to DNA-directed RNA polymerase II largest chain - Mastigamoeba invertens	2e-23	T31670 DNA-directed RNA polymerase (EC 2.7.7.6) II largest chain - Mastig	40.6	12.0	52.9	47.2	3e-85	01 METABOLISM
Pc20g0453 strong similarity to hypothetical protein An01g12010 - Aspergillus niger	5e-19	C68394 protein T24P13.4 [imported] - Arabidopsis thaliana	16.8	12.0	17.3	27.0	8e-82	01 METABOLISM
Pc20g0501 strong similarity to hypothetical 3-hydroxyacyl-CoA-dehydrogenase hcd - Gibberella zeae	7e-34	AB088352 gene: "hcd"; product: "putative 3-hydroxyacyl-CoA-dehydrogenase";	61.7	72.5	99.6	208.6	0	
Pc20g0673 strong similarity to hypothetical homolog of prokaryotic 2-hydroxyhepta-2,4-diene-1,7-diole is	2e-55	T50353 homolog to prokaryotic 2-hydroxyhepta-2,4-diene-1,7-diole isomeras	91.5	108.1	271.1	234.9	1e-111	01.04 phosphate metabolism
Pc20g0875 similarity to hypothetical gap protein - Oryza sativa	2e-10	T118349 probable gap protein - rice blast fungus gypsy retroelement	60.1	100.5	128.2	128.2	0	
Pc20g0955 similarity to hypothetical protein An11g01250 - Aspergillus niger	9e-10	A48665 protein-tyrosine kinase (EC 2.7.1.112) mask, long splice form - huma	12.0	12.0	26.0	51.4	4e-39	08.16 extracellular transport, exocytosis and secretion
Pc20g0985 strong similarity to hypothetical protein An16g07980 - Aspergillus niger	0	0	12.0	12.0	32.3	34.4	0	
Pc20g1005 strong similarity to hypothetical protein SPBC12C2.09c - Schizosaccharomyces pombe	5e-44	AC006733 gene: "Y32H12A.5"; product: "Hypothetical protein Y32H12A.5"; Cai	12.0	12.0	25.8	37.3	1e-76	40.10 nucleus
Pc20g1075 weak similarity to hypothetical protein B210.080 - Neurospora crassa	0	0	78.6	46.6	132.1	164.8	8e-73	01.01.01.07.06 biosynthesis of lysine
Pc20g1105 strong similarity to hypothetical protein contig1414_scaffold.8.1fa_570wg - Aspergillus nid	9e-13	NC80A10_1 gene: "80A10.010"; product: "related to transcriptional activator CMR	12.1	14.3	19.7	31.8	3e-78	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc20g1151 strong similarity to hypothetical protein YOL055w - Saccharomyces cerevisiae	2e-63	NC1074_33 gene: "Y7F4.340"; product: "conserved hypothetical protein"; Neurosp	167.8	158.1	277.2	337.1	0	
Pc20g1265 strong similarity to hypothetical protein BAB09014.1 - Arabidopsis thaliana	1e-93	BX482620 gene: "B11E5.400"; product: "conserved hypothetical protein"; Neur	37.4	36.7	135.6	113.7	0	
Pc20g1273 similarity to hypothetical protein An12g07270 - Aspergillus niger	0	0	12.0	12.0	24.1	44.7	0	
Pc20g1281 similarity to hypothetical protein contig_1_119_scaffold.9.1fa_20cg - Aspergillus nidulans	0	0	19.5	13.4	19.4	27.5	2e-96	01 METABOLISM
Pc20g1281 strong similarity to hypothetical protein contig1485_1.1fa_290cg - Aspergillus fumigatus	0	0	22.7	20.3	42.2	77.8	8e-58	04 TRANSCRIPTION
Pc20g1341 strong similarity to hypothetical protein contig_1_110_scaffold.8.1fa_570wg - Aspergillus nid	5e-55	G72004 8-amino-7-oxononanoate synthase (EC 2.3.1.47) - Chlamydomonas p	137.9	87.8	167.8	229.3	1e-111	01 METABOLISM
Pc20g1351 weak similarity to hypothetical protein An12g03400 - Aspergillus niger	2e-04	AK016708 Mus musculus adult male testis cDNA, RIKEN full-length enriched lib	129.9	103.4	460.5	399.1	0	
Pc20g1391 strong similarity to hypothetical UDP-glucose,sterol transferase gstaA - Aspergillus oryzae	0.0	AB078723 gene: "gstaA"; product: "UDP-glucose,sterol transferase"; Aspergillus	28.4	27.4	53.4	70.7	0	
Pc20g1404 strong similarity to hypothetical mixed-linked glucanase precursor related protein MLG1 - Neu	1e-76	T39920 probable glucanase precursor - fission yeast (Schizosaccharomyces	37.5	28.5	57.7	66.8	6e-51	01.05.04 regulation of C-compound and carbohydrate utilization
Pc20g1421 similarity to hypothetical protein An02g04060 - Aspergillus niger	0	0	277.7	189.3	462.6	461.4	0	
Pc20g1484 similarity to hypothetical monocarboxylate permease Estp6 - Saccharomyces cerevisiae	2e-25	AY236409 gene: "mct"; product: "monocarboxylate transporter-like protein"; Em	77.2	54.3	247.6	255.4	2e-55	01.06.04 breakdown of lipids, fatty acids and isoprenoids
Pc20g1485 strong similarity to hypothetical protein An04g09900 - Aspergillus niger [putative sequencing	0	0	32.7	14.7	43.0	80.1	5e-25	03.01.09.05 DNA conformation modification (e.g. chromatin)
Pc20g1535 strong similarity to hypothetical protein contig1414_scaffold.8.1fa_260wg - Aspergillus fumigatus	7e-20	A71213 alanine-tRNA ligase truncated homolog PH1969 - Pyrococcus horfei	42.8	57.4	95.6	159.3	0	
Pc20g1545 strong similarity to hypothetical protein CAB92029.2 - Neurospora crassa	2e-34	NCB9J10_1 gene: "B9J10.220"; product: "hypothetical protein"; Neurospora cras	134.9	82.8	116.6	177.3	1e-113	01 METABOLISM

Pc20g155: strong similarity to hypothetical protein contig1492_0.1fa_3030cg - Aspergillus fumigatus	4e-62	MUKAD	lysosome (EC 3.2.1.17) - fungus (Chalara sp.)	12.6	12.3	39.0	52.6	0	
Pc20g155: strong similarity to hypothetical protein contig1492_0.1fa_3060wg - Aspergillus fumigatus	1e-150	NCB3E4_1.gene: "B3E4.200"; product: "related to guanine deaminase"; Neurosp		12.0	12.0	27.7	36.1	5e-47	99 UNCLASSIFIED PROTEINS
Pc20g156: strong similarity to hypothetical protein contig_1_100_scaffold_7.1fa_390cg - Aspergillus nidulans	1e-16	AP005945.gene: "blt2936"; Bradyrhizobium japonicum USDA 110 DNA, comple		12.0	12.0	12.0	41.1	9e-84	04.05.01.04 transcriptional control
Pc20g156: strong similarity to hypothetical protein An13g01780 - Aspergillus niger	1e-15	AP004334.gene: "P0455H1.102"; product: "unknown protein"; Oryza sativa (Ja		12.0	12.0	21.1	33.1	6e-76	01.05 C-compound and carbohydrate metabolism
Pc20g157: strong similarity to hypothetical protein mg10764.1 - Magnaporthe grisea	4e-15	S36336	probable transcription factor PLZF - human	37.1	16.4	36.4	36.4	2e-41	01.04.07 phosphate transport
Pc21g0151: strong similarity to hypothetical protein contig31_part.1.1fa_2650cg - Aspergillus fumigatus	4e-38	G72391	conserved hypothetical protein; Thelemonota maritima (strain MS88)	71.0	110.1	145.8	207.1	0	0
Pc21g0161: strong similarity to hypothetical membrane protein YOL119c - Saccharomyces cerevisiae	3e-80	NCB23B1.gene: "B23B10.030"; product: "related to monocarboxylate transport		12.0	12.0	33.7	23.9	0	0
Pc21g023: weak similarity to hypothetical protein T24H10.2 - Caenorhabditis elegans	4e-11	AF361222.gene: "jBA"; product: "JUN-like bZIP transcription factor"; Emericel		524.6	660.9	3263.0	1231.9	0	0
Pc21g028: weak similarity to hypothetical protein An15g05860 - Aspergillus niger				12.0	12.0	31.7	19.7	0	0
Pc21g042: strong similarity to hypothetical protein An14g0630 - Aspergillus niger				10.2	76.6	146.4	105.1	1e-83	99 UNCLASSIFIED PROTEINS
Pc21g049: strong similarity to hypothetical protein CAD2842.1 - Aspergillus fumigatus	2e-69	BX649606	product: "hypothetical protein, conserved"; Aspergillus fumigatus BA	42.1	27.4	51.1	67.5	3e-23	01.01 amino acid metabolism
Pc21g060: similarity to hypothetical membrane domain protein involved in signal transduction AAL22257	3e-11	AE008856.gene: "STM3388"; product: "putative membrane domain protein invol		12.0	12.0	45.3	41.0	0	0
Pc21g061: similarity to hypothetical conserved protein ynaD - Bacillus subtilis	2e-11	NC80A10.gene: "80A10.200"; product: "conserved hypothetical protein"; Neurc		14.4	12.0	52.2	69.7	0	0
Pc21g069: strong similarity to hypothetical protein An02g04420 - Aspergillus niger				0	12.6	12.0	38.9	44.6	0
Pc21g083: similarity to hypothetical protein An14g08230 - Aspergillus niger	5e-11	BX842620.gene: "B11E5.090"; product: "conserved hypothetical protein"; Neurc		12.0	12.1	29.0	27.5	0	0
Pc21g090: hypothetical protein				0	24.7	14.2	47.1	0	0
Pc21g091: strong similarity to hypothetical protein CO0533 - Caulobacter crescentus	1e-108	D87315	conserved hypothetical protein CO0533 [Imported] - Caulobacter cres	184.9	198.2	391.2	488.1	0	0
Pc21g095: weak similarity to hypothetical protein An01g06790 - Aspergillus niger				0	34.5	14.5	35.6	47.1	8e-88
Pc21g098: strong similarity to hypothetical protein contig_1_158_scaffold_13.1fa_200wg - Aspergillus niger		AX196125	Sequence 197 from Patent WO0151639.	24.7	20.5	89.7	72.4	0	0
Pc21g098: strong similarity to hypothetical protein contig1477_1.1fa_1650wg - Aspergillus fumigatus	3e-24	AP005024	product: "hypothetical protein"; Streptomyces avermiltis genomic DN	19.4	22.8	68.0	63.4	0	0
Pc21g104: similarity to hypothetical protein contig_1_46_scaffold_3.1fa_20cg - Aspergillus nidulans				0	13.4	12.0	25.3	37.4	0
Pc21g104: strong similarity to hypothetical protein contig_1_150_scaffold_12.1fa_180cg - Aspergillus niger		AF067182.gene: "idi-2"; product: "IDI-2 precursor"; Podospora anserina IDI-2 pr		0	17.3	22.9	37.7	50.8	5e-65
Pc21g105: strong similarity to hypothetical protein An01g07820 - Aspergillus niger				0	65.3	81.2	119.8	203.3	1e-83
Pc21g108: similarity to MINDBOMB - Homo sapiens	5e-18	AY147849	gene: "MIB"; product: "MINDBOMB"; Homo sapiens MINDBOMB (n	236.8	54.8	542.3	324.2	0	0
Pc21g109: strong similarity to hypothetical regulator protein CAB16735.1 - Schizosaccharomyces pombe	2e-40	T38690	probable regulatory protein - fission yeast (Schizosaccharomyces por	12.0	18.7	48.6	48.8	6e-94	01.05.01 C-compound and carbohydrate utilization
Pc21g109: similarity to hypothetical protein contig_1_26_scaffold_2.1fa_360cg - Aspergillus nidulans				0	15.9	27.5	53.5	70.1	0
Pc21g121: strong similarity to hypothetical oxidoreductase SPAC977.14c - Schizosaccharomyces pombe	1e-116	AY072188.gene: "At1g18270"; product: "unknown protein"; Arabidopsis thaliana		12.0	12.0	13.7	36.1	2e-47	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc21g124: weak similarity to heterokaryon incompatibility protein het-6 - Neurospora crassa				0	12.0	12.0	38.3	49.4	0
Pc21g126: strong similarity to hypothetical protein 1465_scaffold_9.1fa_470wg - Fusarium graminearum	9e-36	C82643	daunorubicin C-13 ketoreductase XF1741 [Imported] - Xylella fastidiosa	585.3	699.2	2024.5	2102.9	0	0
Pc21g130: hypothetical protein				0	12.0	12.7	19.4	37.7	0
Pc21g138: strong similarity to hypothetical protein - Gloeobacter violaceus	4e-30	AP006569	Gloeobacter violaceus PCC 7421 DNA, complete genome, section 2/	19.1	17.3	64.3	79.2	0	0
Pc21g141: strong similarity to hypothetical protein mg06620.1 - Magnaporthe grisea				0	216.3	122.0	181.3	267.7	5e-55
Pc21g145: strong similarity to hypothetical protein contig_1_41_scaffold_2.1fa_40cg - Aspergillus nidulans				0	16.9	15.6	33.6	58.9	0
Pc21g146: strong similarity to hypothetical protein YDR196c - Saccharomyces cerevisiae	5e-42	S52703	hypothetical protein YDR196c - yeast (Saccharomyces cerevisiae)	53.1	62.7	82.4	175.8	1e-50	01.07.01 biosynthesis of vitamins, cofactors, and prosthetic groups
Pc21g146: strong similarity to hypothetical protein An09g02950 - Aspergillus niger	4e-07	T50951	hypothetical protein B24P7.60 [Imported] - Neurospora crassa	23.9	18.0	54.3	59.2	0	0
Pc21g154: weak similarity to hypothetical UbiC COO5 family methyltransferase - Caulobacter crescentus	2e-06	AP005943.gene: "ubiE"; product: "ubiquinone/menaquinone biosynthesis methyl		769.1	614.8	3302.2	1796.0	0	0
Pc21g156: strong similarity to cyanovirin-N like protein An01g05960 - Aspergillus niger	1e-04	NC43E3_4.gene: "43E3.400"; product: "hypothetical protein"; Neurospora crassa		1076.2	1439.8	3096.1	2699.3	0	0
Pc21g158: strong similarity to hypothetical protein contig1495_1.1fa_1490wg - Aspergillus fumigatus				0	37.7	40.9	69.1	83.6	0
Pc21g163: strong similarity to hypothetical protein An09g05240 - Aspergillus niger	2e-38	BX842620.gene: "B11E5.340"; product: "putative protein"; Neurospora crassa C		18.2	14.9	37.0	33.3	0	0
Pc21g168: strong similarity to hypothetical protein ybUJ - Escherichia coli	9e-61	E64819	ybUJ protein - Escherichia coli (strain K-12)	12.4	12.7	35.0	55.0	0	0
Pc21g173: similarity to hypothetical protein SC4B10.22 - Streptomyces collicolor	1e-13	SC039315.gene: "SC07121"; "SC4B10.22"; product: "putative secreted protein		12.0	12.3	23.7	52.7	2e-90	01.05.01 C-compound and carbohydrate utilization
Pc21g182: weak similarity to hypothetical protein An17g00620 - Aspergillus niger				0	14.4	12.0	14.8	31.1	0
Pc21g186: strong similarity to hypothetical protein mg06182.1 - Magnaporthe grisea	1e-96	T52133	potassium channel beta subunit homolog [Imported] - Arabidopsis th	227.5	268.8	477.4	618.9	2e-68	01 METABOLISM
Pc21g186: similarity to hypothetical protein CC3654 - Caulobacter crescentus	5e-18	AP005943.gene: "blt2446"; Bradyrhizobium japonicum USDA 110 DNA, comple		12.0	12.0	41.6	55.5	0	0
Pc21g197: strong similarity to hypothetical protein contig31_part.1.1fa_3670wg - Aspergillus fumigatus	1e-09	AD1155	conserved hypothetical protein homolog lmo644 [Imported] - Listeri	79.3	98.3	160.0	155.7	1e-111	01.03.16 polynucleotide degradation
Pc21g200: strong similarity to hypothetical methyltransferase AAO34671.1 - Gibberella zeae	1e-39	AF39361.gene: "OrfF"; product: "methyltransferase"; Gibberella zeae strain G2		14.2	12.0	43.3	65.7	0	0
Pc21g201: strong similarity to hypothetical protein contig1492_0.1fa_3170cg - Aspergillus fumigatus	7e-92	NCB11625.gene: "B11B23.050"; product: "related to lariat-debranching enzyme"		31.2	23.0	55.0	80.9	0	0
Pc21g214: similarity to hypothetical protein An09g03770 - Aspergillus niger				0	12.0	12.0	41.9	0	0
Pc21g214: hypothetical protein				0	27.9	22.1	90.9	55.5	0
Pc21g218: hypothetical protein				0	12.0	12.0	14.4	36.1	0
Pc21g219: weak similarity to hypothetical protein contig_1_149_scaffold_12.1fa_40cg - Aspergillus nidulans				0	19.8	14.1	19.5	53.1	5e-38
Pc21g221: strong similarity to hypothetical protein 1316_scaffold_4.1fa_360cg - Fusarium graminearum	5e-08	AY057845	product: "unknown"; Zymomonas mobilis strain ZM4 plasmid 1, com	141.5	44.6	55.5	184.3	0	0
Pc21g228: strong similarity to transcription regulator alcR - Aspergillus nidulans	0.0	ALCR	EM REGULATORY PROTEIN ALCR	108.0	112.8	142.8	250.9	1e-86	08.16.03 Type I protein secretion system (ABC-type transport systems)
Pc21g235: weak similarity to hypothetical glutathione S-transferase BA06940.1 - Gibberella fujikuroi	8e-07	AB071861	product: "putative glutathione S-transferase"; Gibberella fujikuroi mR1	16.0	13.0	53.3	38.1	0	0
Pc21g236: strong similarity to hypothetical protein contig46_part.1.1fa_2310cg - Aspergillus fumigatus	7e-66	BX897679	gene: "B2C2.090"; product: "probable positive effector protein GCN	46.9	89.5	224.5	150.3	3e-21	03.01.99 other DNA processing
Pc21g235: strong similarity to hypothetical protein CAD2381.1 - Neurospora crassa	8e-28	NC12344.gene: "123A4.300"; product: "conserved hypothetical protein"; Neurc		257.9	352.7	572.3	1016.6	0	0
Pc21g236: similarity to protease synthase and sporulation negative regulatory protein pai1 - Enterococcus faecalis	1e-18	AE016956	product: "protease synthase and sporulation negative regulatory protein	12.6	12.0	26.5	41.0	0	0
Pc22g004: weak similarity to hypothetical protein T1G12.10 - Arabidopsis thaliana	1e-06	T40752	hypothetical protein SPBC8D2.07c - fission yeast (Schizosaccharom	37.0	36.7	70.8	91.4	0	0
Pc22g005: strong similarity to hypothetical conserved protein PA2682 - Pseudomonas aeruginosa	1e-56	D83310	conserved hypothetical protein PA2682 [Imported] - Pseudomonas ae	39.8	73.5	129.0	169.4	1e-170	10 CELLULAR COMMUNICATION/SIGNAL TRANSDUCTION MECHANISM
Pc22g006: strong similarity to hypothetical protein CO0533 - Caulobacter crescentus	1e-95	D87315	conserved hypothetical protein CO0533 [Imported] - Caulobacter cres	12.8	12.0	28.6	39.1	0	0
Pc22g016: weak similarity to arkyrin AAC37208.1 - Drosophila melanogaster	1e-28	T13940	ar-kyrin - fruit fly (Drosophila melanogaster)	51.1	32.0	148.4	140.0	7e-72	01.02.99 other nitrogen and sulfur metabolism activities
Pc22g019: similarity to hypothetical protein contig1492_0.1fa_5060wg - Aspergillus fumigatus				0	226.6	199.7	440.8	530.4	0
Pc22g034: strong similarity to hypothetical protein aq_928 - Aquifex aeolicus	3e-67	G90073	conserved hypothetical protein SA2448 [Imported] - Staphylococcus i	19.9	12.0	21.5	35.7	1e-71	03.03.01 mitotic cell cycle and cell cycle control
Pc22g046: strong similarity to hypothetical protein contig12.1fa_1670wg - Aspergillus fumigatus				0	54.7	84.6	112.7	111.6	0
Pc22g062: similarity to hypothetical AAA-ATPase AAR34299.1 - Geobacter sulfurreducens	3e-16	AE017210	product: "ATPase, AAA family"; Geobacter sulfurreducens PCA, sex	12.0	12.0	31.3	40.1	0	0
Pc22g065: similarity to hypothetical protein 1193_scaffold_2.1fa_450wg - Fusarium graminearum	4e-07	T45428	hypothetical protein MLCB37.35c [Imported] - Mycobacterium lepra	12.0	12.0	36.6	58.7	0	0
Pc22g068: strong similarity to hypothetical protein ydcF - Escherichia coli	1e-38	E85737	hypothetical protein ydcF [Imported] - Escherichia coli (strain O157_H	45.6	25.1	54.2	57.5	0	0
Pc22g074: strong similarity to hypothetical protein An11g09420 - Aspergillus niger				0	12.0	12.0	32.5	38.3	0
Pc22g075: similarity to hypothetical protein contig202.1fa_130cg - Aspergillus fumigatus				0	98.1	75.0	146.2	206.7	2e-21
Pc22g076: similarity to hypothetical protein DRB0099 - Deinococcus radiodurans	2e-16	A75629	hypothetical protein - Deinococcus radiodurans (strain R1)	15.5	12.0	36.8	50.9	2e-51	01 METABOLISM
Pc22g076: weak similarity to hypothetical thiosulfate sulfurtransferase - Pseudomonas aeruginosa	1e-125	F83319	probable thiosulfate sulfurtransferase PA2603 [Imported] - Pseudom	60.2	94.3	120.0	262.3	0	0
Pc22g082: strong similarity to hypothetical protein - Danio rerio	2e-33	BC066536	product: "Unknown (protein for MGC:76849)"; Danio rerio cDNA clon	12.6	12.0	44.9	50.6	0	0
Pc22g082: hypothetical protein				0	25.3	16.9	47.0	34.6	0
Pc22g088: strong similarity to hypothetical protein An02g14010 - Aspergillus niger				0	37.3	37.8	135.2	97.5	1e-141
Pc22g089: weak similarity to hypothetical protein alf7165 - Nostoc sp.	3e-04	F96014	conserved hypothetical protein SMb20675 [Imported] - Sinorhizobium	84.5	133.3	317.4	403.3	2e-57	01 METABOLISM
Pc22g094: strong similarity to hypothetical protein T16K5.230 - Arabidopsis thaliana	8e-39	AE016941	product: "conserved hypothetical protein"; Bacteroides thetaiotaomic	12.0	12.0	12.0	26.4	0	0
Pc22g128: similarity to hypothetical protein F24K9.9 - Arabidopsis thaliana	2e-09	T02539	hypothetical protein At2g37730 [Imported] - Arabidopsis thaliana	159.2	159.3	197.9	385.3	1e-104	03.01.03 DNA synthesis and replication
Pc22g135: hypothetical protein				0	12.0	12.0	12.0	29.0	4e-23
Pc22g140: strong similarity to hypothetical double strand break catalysing ATSP011.1 - Arabidopsis thaliana	2e-37	T48781	MEIOTIC RECOMBINATION PROTEIN REC12 related protein [Imp	12.0	12.0	40.6	30.1	2e-84	01.02.01 nitrogen and sulfur utilization
Pc22g141: strong similarity to hypothetical membrane protein YOL092w - Saccharomyces cerevisiae	1e-48	S53737	probable membrane protein YOL092w - yeast (Saccharomyces cerev	47.5	37.9	63.7	76.0	0	0
Pc22g142: strong similarity to hypothetical protein contig5_part.1.1fa_390wg - Aspergillus fumigatus	4e-99	BX572599	product: "possible 2-nitropropane dioxygenase"; Rhodospseudomon	231.2	118.9	318.6	333.5	1e-102	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc22g145: strong similarity to hypothetical protein An07g01270 - Aspergillus niger				0	12.0	19.8	34.7	32.2	0.0
Pc22g156: similarity to hypothetical membrane protein YPR147c - yeast (Saccharomyces cerevisiae)	6e-21	S69034	probable membrane protein YPR147c - yeast (Saccharomyces cerev	15.2	12.0	24.7	30.7	1e-19	04.05.01.04 transcriptional control
Pc22g166: strong similarity to hypothetical membrane protein YL067c - Saccharomyces cerevisiae	1e-121	NC5F3_22.gene: "SF3.220"; product: "conserved hypothetical protein"; Neurosp		56.5	65.2	91.9	163.1	0	0
Pc22g172: strong similarity to hypothetical protein 1185_scaffold_2.1fa_260wg - Fusarium graminearum	6e-11	AP002994.gene: "mib0013"; product: "transcriptional regulator"; Mesorhizobium		13.2	23.1	37.0	61.8	0	0
Pc22g177: strong similarity to hypothetical protein An11g02090 - Aspergillus niger	1e-35	NCB19A11.gene: "B19A117.170"; product: "hypothetical protein"; Neurospora cr		22.4	12.0	47.0	39.6	0	0
Pc22g195: weak similarity to hypothetical protein An07g04490 - Aspergillus niger				0	46.9	74.7	94.4	171.3	1e-70
Pc22g201: weak similarity to hypothetical protein An13g02200 - Aspergillus niger				0	20.8	19.3	26.6	42.3	0
Pc22g205: weak similarity to hypothetical RNA polymerase II transcription factor Upc2 - Saccharomyces cerevisiae	4e-05	S51451	probable membrane protein YLR228c - yeast (Saccharomyces cerev	18.2	20.2	41.4	75.6	1e-83	01.06.04 breakdown of lipids, fatty acids and isoprenoids
Pc22g207: strong similarity to hypothetical protein An14g03100 - Aspergillus niger				0	26.6	23.5	52.4	61.7	1e-125
Pc22g210: strong similarity to hypothetical protein contig31_part_1.1fa_2380cg - Aspergillus fumigatus	[6e-13]	AE016891	Chromobacterium violaceum ATCC 12472 section 2 of 16 of the	16.8	23.4	24.2	70.1	0	0
Pc22g214: strong similarity to hypothetical protein mg06547.1 - Magnaporthe grisea	1e-142	AV32375.gene: "ttuC"; product: "tartrate dehydrogenase"; Agrobacterium vitis		47.8	36.4	45.9	82.1	0	0

Pc22g216c similarity to hypothetical protein contig_1_18_scaffold_1.tfa_110wg - Aspergillus nidulans	0	0	0	22.1	12.0	47.9	51.6	0		0
Pc22g217c strong similarity to hypothetical protein contig1490_3.tfa_1220cg - Aspergillus fumigatus	8e-24	NCB13H1f gene: "B13H18.210"; product: "related to G protein coupled receptor	0	30.7	33.4	61.7	112.0	1e-29	99 UNCLASSIFIED PROTEINS	0
Pc22g223i hypothetical protein	0	0	0	12.0	12.0	24.9	32.8	0		0
Pc22g230c strong similarity to hypothetical protein An12g07700 - Aspergillus niger	5e-07	AE015943 gene: "CTC02163"; product: "ethanolamine utilization protein (transc	0	16.3	16.9	31.3	46.5	0		0
Pc22g231f strong similarity to hypothetical protein An03g06340 - Aspergillus niger	0	0	0	12.0	12.0	21.6	43.0	0		0
Pc22g232i strong similarity to hypothetical protein An03g06370 - Aspergillus niger	0	0	0	12.0	12.0	16.8	28.8	1e-64	03.03.01 mitotic cell cycle and cell cycle control	0
Pc22g239c strong similarity to hypothetical protein ncu07110.1 - Neurospora crassa	0	0	0	25.0	33.5	53.9	66.7	0		0
Pc22g241f strong similarity to hypothetical protein contig42.tfa_50wg - Aspergillus fumigatus	4e-65	AP006575 Gloeobacter violaceus PCC 7421 DNA, complete genome, section 8/	0	20.1	26.0	43.9	58.8	3e-39	01 METABOLISM	0
Pc22g241i strong similarity to hypothetical protein An18g01930 - Aspergillus niger	0	0	0	14.2	19.0	36.4	28.5	0		0
Pc22g242c weak similarity to hypothetical isochorismatase Tso729 - Thermoplasma acidophilum	1e-22	AE016863 product: "isochorismatase family protein"; Pseudomonas syringae pv	0	67.3	75.6	201.6	242.6	0		0
Pc22g249f weak similarity to hypothetical protein AAM35689.1 - Xanthomonas axonopodis	3e-13	BX649607 product: "hypothetical protein, conserved"; Aspergillus fumigatus BA	0	12.0	12.0	26.5	30.9	0		0
Pc22g252c similarity to hypothetical protein An18g06380 - Aspergillus niger	0	0	0	13.6	12.0	34.3	41.6	0		0
Pc22g255c strong similarity to hypothetical protein contig1497_2.tfa_50wg - Aspergillus fumigatus	0	0	0	484.2	567.4	1130.3	1145.9	0		0
Pc23g005c strong similarity to hypothetical protein An07g04950 - Aspergillus niger [putative sequencing 4e-06	4e-06	AY258009 product: "Orf17"; Streptomyces clavuligerus clavulanic acid biosynth	0	12.0	12.0	39.4	20.1	0		0

Supplementary Table 14. K-mean cluster 6

⑧=Values given are the average of three independent experiments

orf code	Description of putative <i>P. chrysogenum</i> ORF	Best blast homologs to putative <i>P. chrysogenum</i> ORF		Average transcript levels in		Function (auto)	Category
		[e-value]	Gene code	WIS - PAA	WIS + PAA	DS - PAA	
Pc10g0070	weak similarity to germinase 1.2-dioxygenases GDO - <i>Pseudomonas alcaligenes</i>	0.0	BX68692_119	19.8	24.3	134.1	01.05.01 C-compound and carbohydrate utilization
Pc12g01540	strong similarity to sulfite permease SubB - <i>Penicillium chrysogenum</i>	0.0	AF163974_1	11.1	12.0	33.6	01.05.01 C-compound and carbohydrate utilization
Pc12g04340	strong similarity to precursor of alpha-amylase A - <i>Aspergillus niger</i>	6e-96	AT0AAG2_1	15.27	12.0	106.7	01.05.01 C-compound and carbohydrate utilization
Pc12g07100	strong similarity to alcohol dehydrogenase adhB1 - <i>Aspergillus nidulans</i>	1e-107	ADH10228_1	24.1	12.0	142.6	01.05.01.01 sugar, glucoside, polyol and carboxylate catabolism
Pc12g12910	strong similarity to endo-alpha-1,5-arabinanase abnA - <i>Aspergillus niger</i>	4e-04	AZ7881_1	49.9	23.7	48.8	01.05.07 C-compound, carbohydrate transport
Pc13g04530	weak similarity to cell wall synthesis protein KRE9 - <i>Candida albicans</i>	6e-52	AF046252_1	47.4	12.0	123.0	01.03 nucleotide metabolism
Pc13g07200	strong similarity to carboxylic acid transport protein Jen1 - <i>Saccharomyces cerevisiae</i>	0.0	AF002028_1	61.0	14.3	102.6	01.05.07 C-compound, carbohydrate transport
Pc13g07630	strong similarity to MIP dehydrogenase IMH3 - <i>Candida albicans</i>	0.0	BMH_CANAL	556.6	48.4	206.9	01.03.04 pyrimidine nucleotide metabolism
Pc13g13280	strong similarity to carnitine acetyl transferase hscC - <i>Aspergillus nidulans</i>	1e-131	BX68692_112	262.8	181.5	523.9	01.03.04 nucleotide metabolism
Pc13g13450	strong similarity to purine-cytosine permease pcyD - <i>Kluyveromyces marxianus</i>	8e-61	XM0011419_1	27.6	12.0	33.5	03.01.03 DNA synthesis and replication
Pc16g03600	strong similarity to phosphotriester acetyltransferase like protein An04G0830 - <i>Aspergillus niger</i>	2e-08	NCB1383_5	42.6	24.0	46.7	01.01.01.15.03 biosynthesis of leucine
Pc16g04000	strong similarity to DNA polymerase lambdaB, POLD, lambdaB - <i>Homo sapiens</i> [putative pseudogene]	9e-64	BC025557_1	15.0	12.0	56.2	01.01.01.01 C-compound, carbohydrate utilization
Pc16g07660	strong similarity to long-chain acyl-CoA dehydrogenase like protein An02G0230 - <i>Aspergillus niger</i>	1e-178	AY033336_1	375.9	155.2	149.1	01.05.01.01 C-compound, carbohydrate catabolism
Pc16g10900	strong similarity to beta-ketothiolase dehydrogenase precursor suc1 - <i>Aspergillus niger</i>	0.0	ASY28046_1	12.0	12.0	23.6	01.05.01 C-compound and carbohydrate utilization
Pc20g01070	strong similarity to beta-glucosidase 1 bg11 - <i>Aspergillus aculeatus</i>	0.0	TR05680_1	91.7	35.6	69.0	01.05.07 C-compound, carbohydrate transport
Pc20g14200	strong similarity to beta-glucan synthase associated protein KneB - <i>Saccharomyces cerevisiae</i>	1e-144	AY371202_1	14.4	12.0	27.2	01.05.07 C-compound, carbohydrate transport
Pc20g15720	strong similarity to high affinity hexose transporter Hxt1 - <i>Saccharomyces cerevisiae</i>	1e-89	NC13E11_7	46.8	22.9	39.7	01.01.10.05.02 degradation of cysteine
Pc20g19210	similarly to hypothetical amine transporter SPOC18.02 - <i>Schizosaccharomyces pombe</i>	2e-22	T4145	192.1	28.1	95.6	01.05.07 C-compound, carbohydrate transport
Pc21g04760	strong similarity to cytosine dioxygenase CDO1 - <i>Rattus norvegicus</i>	2e-29	BC020414_1	109.5	23.0	157.7	01.05.07 C-compound, carbohydrate transport
Pc21g10870	strong similarity to fatty-acyl-CoA synthase beta chain fat p1 - <i>Schizosaccharomyces pombe</i>	0.0	S37178	202.5	106.7	281.6	01.05.04 regulation of C-compound and carbohydrate utilization
Pc21g12820	strong similarity to 3-dehydroshikimate dehydrogenase qh-4 - <i>Neurospora crassa</i>	1e-89	BX68692_32	223.3	63.4	943.2	01.05.04 regulation of C-compound and carbohydrate utilization
Pc21g12830	strong similarity to hypothetical amine transporter SPOC18.02 - <i>Schizosaccharomyces pombe</i>	9e-38	T4145	83.1	20.6	156.0	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc21g15430	strong similarity to dimethylallyltransferase synthase dmwA1 - <i>Clostridium purpureum</i>	2e-56	AF259382_1	883.9	299.6	923.6	01.20.37 C-biosynthesis of peptide antibiotics
Pc21g15470	similarly to mitochondrial 25-hydroxyvitamin D3 24-hydroxylase CYP50C24 - <i>Gallus gallus</i>	6e-15	AL138806_1	24.7	11.5	74.2	01.04 phosphate metabolism
Pc21g15480	strong similarity to cyclic peptide AM-toxin synthase like protein An02G02650 - <i>Aspergillus niger</i>	0.0	MAPEPPTNT_1	562.2	28.9	449.2	01.05.01 C-compound and carbohydrate utilization
Pc21g15790	strong similarity to chitinase precursor CH33 - <i>Trichoderma reesei</i> [putative sequencing error]	1e-104	AF510301_1	26.8	10.0	41.3	01.05.01 C-compound and carbohydrate utilization
Pc21g16210	strong similarity to scytalone dehydratase - <i>Piculicaria cryzae</i>	8e-62	U95042_1	80.2	39.2	43.0	01.04 phosphate metabolism
Pc21g22620	strong similarity to alcohol dehydrogenase actA - <i>Aspergillus nidulans</i>	1e-169	A25654	181.5	71.8	313.9	01.05.01 C-compound, carbohydrate catabolism
Pc21g25200	strong similarity to extracellular alpha-glucosidase xgl1 - <i>Aspergillus niger</i>	0.0	NC04611_13	221.4	44.3	131.2	01.05.01 C-compound and carbohydrate utilization
Pc21g25450	strong similarity to xylosidase xnbD - <i>Aspergillus niger</i>	0.0	JC7966	120.0	12.0	120.0	01.05.01 C-compound and carbohydrate utilization
Pc22g01100	strong similarity to chitinase 1 precursor cdt1 - <i>Coccidioides immitis</i>	1e-158	AY271350_1	43.0	59.9	300.4	01.05.07 C-compound, carbohydrate transport
Pc22g10150	glucan 1,4-glucanase BAC0261 - <i>Aspergillus fumigatus</i>	0.0	AB091510_1	18.6	12.0	31.6	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc22g14640	strong similarity to high affinity glucose transporter HGT1 - <i>Kluyveromyces fragilis</i>	1e-130	AB186113_2	38.1	12.0	31.5	01.05.07 C-compound, carbohydrate transport
Pc22g25490	weak similarity to 2-haloacid halohydratase IvaA - <i>Burkholderia cepacia</i>	9e-36	NCB1182_26	59.3	15.0	50.6	01.05.07 C-compound, carbohydrate transport
Pc22g27130	strong similarity to 4-hydroxyphenylpyruvate decarboxase HPPD - <i>Mycophtharella graminicola</i>	1e-124	4-HPPD_MYCCOR	192.4	106.1	316.6	01.05.07 C-compound and carbohydrate metabolism
Pc22g27470	strong similarity to hexose transporter Hxt13 - <i>Saccharomyces cerevisiae</i>	9e-82	NC13E11_7	14.3	12.0	22.6	01.05.07 C-compound and carbohydrate metabolism
Pc22g28420	strong similarity to brown 2 protein abt2 - <i>Aspergillus fumigatus</i>	0.0	AF104823_1	34.0	12.0	62.1	01.02.01.09.98 other catabolism of nitrogenous compounds
Pc22g13330	strong similarity to glycerol kinase GK - <i>Mus musculus</i>	1e-139	AF076623_1	39.5	20.7	121.9	01.03.02 nucleic acid
Pc22g16830	strong similarity to lactone-specific esterase estI1 - <i>Pseudomonas fluorescens</i>	2e-71	F83383	91.8	34.2	36.1	01.02.01 nitrogen and sulfur utilization
Pc22g19620	weak similarity to 35.6K acid phosphatase - <i>Aspergillus nidulans</i>	3e-36	NCB21212_12	86.1	26.9	228.3	01.05.07 C-compound and carbohydrate metabolism
Pc22g20960	strong similarity to protein kinase Htt1 - <i>Aspergillus nidulans</i>	1e-143	A33697	10.0	12.0	13.6	01.05.07 C-compound and carbohydrate metabolism
Pc22g22650	strong similarity to androgen-inducible aldehyde reductase alar - <i>Rattus norvegicus</i>	3e-32	HS4271799_1	48.1	36.4	151.4	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc22g22820	strong similarity to lipase LipP - <i>Pseudomonas sp.</i>	0.0	AF034086_1	120.3	110.1	163.9	01.05.07 C-compound and carbohydrate metabolism
Pc22g24730	strong similarity to delta12 fatty acid desaturase - <i>Aspergillus niger</i>	1e-63	AB066061_129	59.9	12.0	17.4	01.05.07 C-compound and carbohydrate metabolism
Pc22g24830	strong similarity to carboxyphosphonohydrolyate phosphonumutase bcpA - <i>Streptomyces hygroscopicus</i>	1e-155	AN0567910_1	56.6	13.6	12.6	01.04 phosphate metabolism
Pc10g00010	weak similarity to kinesin-related protein KLP4 - <i>Aspergillus nidulans</i>	0	0	292.0	100.0	539.6	01.05.07 C-compound and carbohydrate metabolism
Pc22g23660	strong similarity to Swei1 regulating protein kinase Htt1 - <i>Saccharomyces cerevisiae</i>	3e-93	KLAE55232_1	36.1	10.0	12.0	01.05.01.04 transcriptional control
Pc14g11250	strong similarity to gene expression regulator like protein An12G0080 - <i>Aspergillus niger</i>	0.0	A324	24.4	23.1	63.2	01.04 regulation of amino acid metabolism
Pc16g08010	strong similarity to protein abnA - <i>Aspergillus nidulans</i>	0.0	C326	22.6	12.0	63.5	01.05.07 C-compound and carbohydrate utilization
Pc16g15320	strong similarity to cutinase transcription factor beta CTF1b - <i>Fusarium solii</i>	3e-42	CT1b_FUSSO	23.6	62.9	62.9	06.01 protein binding and stabilization
Pc22g0240	weak similarity to hypothetical transcription regulator protein - <i>Schizosaccharomyces pombe</i>	2e-14	BX68622_15	21.3	23.9	63.5	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc12g04400	strong similarity to heat-shock protein hsp30 - <i>Aspergillus nidulans</i>	1e-25	AB128988_1	87.7	58.5	139.2	01.05.07 C-compound and carbohydrate utilization
Pc12g15740	similarly to lysosomal protease CLN2 - <i>Rattus norvegicus</i>	1e-159	AB046899_1	86.3	59.5	201.2	01.05.07 C-compound and carbohydrate utilization
Pc21g11160	alkaline serine protease AM03367 - <i>Penicillium chrysogenum</i>	1e-175	AF321109_1	74.4	51.0	147.4	01.05.07 C-compound and carbohydrate utilization
Pc21g27400	strong similarity to vacuolar H ⁺ Ca2+/exchanger Vcx1 - <i>Saccharomyces cerevisiae</i>	0.0	T4141	154.6	170.1	250.2	01.05.07 C-compound and carbohydrate utilization
Pc21g27570	strong similarity to vacuolar H ⁺ Ca2+/exchanger Vcx1 - <i>Saccharomyces cerevisiae</i>	1e-39	AF053229_1	245.6	185.8	191.2	01.05.07 C-compound and carbohydrate utilization
Pc21g2850	weak similarity to ecto-ATPase sciam105 - <i>Rattus norvegicus</i>	0	0	29.1	12.0	61.6	01.05.07 C-compound and carbohydrate utilization
Pc21g16380	strong similarity to cell surface fenoxidase precursor Fe3 - <i>Saccharomyces cerevisiae</i>	0.0	AF116091_1	31.3	12.0	61.5	03.03.01 mitotic cell cycle and cell cycle control
Pc22g06520	strong similarity to potassium transporter protein Tk2 - <i>Saccharomyces cerevisiae</i>	2e-47	SC0206453_1	12.0	12.0	33.7	01.05.07 C-compound and carbohydrate utilization
Pc21g07370	strong similarity to negative transcription factor hnt1 - <i>Podospira anserina</i> [putative sequencing error]	1e-158	AF323561_1	23.1	12.0	61.5	01.05.07 C-compound and carbohydrate utilization
Pc12g05710	weak similarity to chemotactic receptor CAR1 - <i>Dicystotium discoidum</i>	2e-69	NCB13148_19	12.3	12.0	47.0	01.05.07 C-compound and carbohydrate utilization
Pc21g7300	weak similarity to cytochrome-N-CV-N - <i>Nostoc ellipsosporum</i>	2e-12	NC43163_1	28.6	13.0	25.0	01.05.07 C-compound and carbohydrate utilization
Pc21g26290	strong similarity to eukaryotic protein precursor	1e-32	AN07412_1	34.5	12.0	61.5	01.07 detoxification
Pc13g03440	strong similarity to capsular associated protein CAP10 - <i>Filobacteriella neofumens</i>	1e-47	AF144574_1	29.9	31.2	58.3	01.05.07 C-compound and carbohydrate utilization
Pc13g12500	weak similarity to ribogalactanase resistance factor Sng1 - <i>Saccharomyces cerevisiae</i>	7e-07	S53820	12.0	12.0	42.0	01.05.07 C-compound and carbohydrate utilization
Pc13g14910	strong similarity to copper resistance-associated P-type ATPase CRP1 - <i>Candida albicans</i>	0.0	AF110558_1	19.1	12.0	44.1	01.05.07 C-compound and carbohydrate utilization
Pc19g02820	strong similarity to cytidine hydratase Cht - <i>Gloeosporium sorghi</i>	1e-140	AY259111_1	12.0	12.0	28.3	01.07 detoxification
Pc20g1760	strong similarity to ceratoprotein transporter CFP - <i>Cercospora kikuchi</i>	1e-137	AF236222_1	44.8	12.0	42.6	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc20g14340	strong similarity to hypothetical 7-aminocycloheptanone resistance protein RTA1 - <i>Aspergillus nidulans</i>	2e-21	S34636	17.4	27.6	31.6	01.05.04 regulation of C-compound and carbohydrate utilization
Pc19g0770	strong similarity to pisinin demethylase PDA6-1 - <i>Nectria haematococca</i>	2e-55	S34396	144.9	52.0	208.9	01.05.04 regulation of C-compound and carbohydrate utilization
Pc19g08010	strong similarity to hypothetical membrane transport protein SPAC31.06 - <i>Schizosaccharomyces pombe</i>	3e-32	AF183691_1	12.8	12.0	38.3	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc21g15240	strong similarity to ceratoprotein transporter CFP - <i>Cercospora kikuchi</i>	1e-153	AF236222_1	44.8	12.0	42.6	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc12g15330	strong similarity to hypothetical cytochrome P450 monooxygenase p450-4 - <i>Gibberella fujikuroi</i>	1e-105	AF278602_3	28.4	15.5	113.9	01.11 electron transport and membrane-associated energy conservation
Pc22g24400	similarly to ferriic cupric reductase Fre2 - <i>Saccharomyces cerevisiae</i>	0.0	BX68622_23	80.48	265.1	508.0	01.03 nucleotide metabolism
Pc12g10870	similarly to ankryin 3 (spice form 3) - <i>Mus musculus</i>	3e-31	T42714	34.0	24.0	55.6	01.05.07 C-compound and carbohydrate utilization
Pc20g0210	strong similarity to secretory component ECF - <i>Saccharophaga peregrina</i>	1e-138	NC5753_11	41.4	11.7	100.6	01.05.07 C-compound and carbohydrate utilization
Pc06g01530	similarly to hypothetical protein 4 - <i>Streptomyces coelicolor</i>	2e-30	AC117176_51	33.9	24.4	56.0	01.05.07 C-compound and carbohydrate utilization
Pc06g02040	similarly to hypothetical protein 15_scaffold_12_1b_520g - <i>Fusarium graminearum</i>	0	0	276.5	144.0	622.6	01.05.07 C-compound and carbohydrate utilization
Pc06g02050	strong similarity to hypothetical protein An01G1190 - <i>Aspergillus niger</i>	0	0	97.3	23.5	61.6	01.05.07 C-compound and carbohydrate utilization
Pc06g02070	weak similarity to hypothetical protein contig946_part_11b_1990g - <i>Aspergillus fumigatus</i>	0	0	12.0	12.0	50.6	01.05.07 C-compound and carbohydrate utilization
Pc06g02070	weak similarity to hypothetical protein contig946_part_11b_1990g - <i>Aspergillus fumigatus</i>	0	0	39.9	12.0	50.6	01.05.07 C-compound and carbohydrate utilization
Pc06g02080	weak similarity to hypothetical protein An02G0230 - <i>Aspergillus niger</i>	0	0	12.0	12.0	61.5	01.05.07 C-compound and carbohydrate utilization
Pc06g02080	weak similarity to hypothetical protein An02G0230 - <i>Aspergillus niger</i>	0	0	12.0	12.0	61.5	01.05.07 C-compound and carbohydrate utilization
Pc06g02090	hypothetical protein	0	0	12.0	12.0	61.5	01.05.07 C-compound and carbohydrate utilization
Pc06g02100	hypothetical protein	0	0	12.0	12.0	61.5	01.05.07 C-compound and carbohydrate utilization
Pc06g02160	hypothetical protein	0	0	866.8	245.0	579.3	03.01.05.01 DNA repair
Pc22g27220	weak similarity to NAD ⁺ -ADP-ribosyltransferase PPOL - <i>Homo sapiens</i>	0.0	AF345753_1	15.7	12.0	25.5	01.03 nucleotide metabolism
Pc22g2850	weak similarity to hypothetical protein - <i>Aspergillus fumigatus</i>	7e-49	BX68692_42	30.3	12.0	57.7	01.05.07 C-compound and carbohydrate utilization
Pc22g2930	strong similarity to oxidoreductase like protein An02G1260 - <i>Aspergillus niger</i>	2e-18	AE03435_30	12.0	12.0	24.5	01.05.07 C-compound and carbohydrate utilization
Pc12g03260	strong similarity to hypothetical protein An02G0730 - <i>Aspergillus niger</i>	0	0	61.5	16.1	147.2	01.05.07 C-compound and carbohydrate utilization
Pc12g04540	strong similarity to hypothetical protein contig_1_55_scaffold_31b_110g - <i>Aspergillus nidulans</i>	0	0	12.0	12.0	29.6	01.05.07 C-compound and carbohydrate utilization
Pc12g06460	strong similarity to hypothetical protein contig281b_310g - <i>Aspergillus fumigatus</i>	0	0	12.0	12.0	123.8	01.04 phosphate metabolism
Pc12g06880	hypothetical protein	0	0	117.0	24.4	288.0	01.05.07 C-compound and carbohydrate utilization
Pc12g06900	strong similarity to hypothetical protein An01G1790 - <i>Aspergillus niger</i>	0	0	24.4	12.0	47.7	03.03.01.01.11 mitosis
Pc12g07740	similarly to hypothetical protein contig_1_128_scaffold_101b_800g - <i>Aspergillus nidulans</i>	0	0	86.4	35.3	92.2	01.05.07 C-compound and carbohydrate utilization
Pc12g09800	strong similarity to hypothetical protein SPAC837.13c - <i>Schizosaccharomyces pombe</i>	1e-40	T39005	140.5	168.8	219.7	01.05.07 C-compound and carbohydrate utilization
Pc12g10150	strong similarity to hypothetical protein An01G1790 - <i>Aspergillus niger</i>	0	0	18.4	39.0	33.4	01.05.07 C-compound and carbohydrate utilization
Pc12g12350	hypothetical protein	0	0	46.9	31.9	58.6	01.05.01.01.02 polysaccharide degradation
Pc12g13160	strong similarity to hypothetical mixed-linked glucanase precursor related protein MLG1 - <i>Neurospora crassa</i>	4e-69	U8106_1	84.0	32.7	67.1	01.05.07 C-compound and carbohydrate utilization
Pc12g13430	strong similarity to hypothetical protein B23.23_120 - <i>Neurospora crassa</i>	1e-132	BC2323_12	12.0	12.0	39.0	01.05.07 C-compound and carbohydrate utilization
Pc12g13840	strong similarity to hypothetical protein contig_1_41_scaffold_21b_105g - <i>Aspergillus nidulans</i>	0	0	12.0	12.0	57.7	01.05.07 C-compound and carbohydrate utilization
Pc12g14130	similarly to hypothetical protein An03G0230 - <i>Agrobacterium tumefaciens</i>	3e-14	AF005951_40	16.1	12.0	105.6	01.05.07 C-compound and carbohydrate utilization
Pc12g14880	strong similarity to hypothetical protein An02G0730 - <i>Aspergillus niger</i>	8e-06	BX68692_119	141.5	17.6	126.4	01.05.07 C-compound and carbohydrate utilization
Pc12g15480	strong similarity to hypothetical protein contig_1_51_scaffold_61b_720g - <i>Aspergillus nidulans</i>	0.0	AF002461_63	12.0	12.0	25.0	01.05.07 C-compound and carbohydrate utilization
Pc12g16600							

Pc13g09460	similarly to zonadhesin - Mus musculus	2e-18	T42215	zonadhesin - mouse	772.1	730.7	2458.2	646.9	0	
Pc13g1090	hypothetical protein	0	0	120.0	120.0	284	12.0	0		
Pc13g13860	strong similarity to hypothetical protein contig_1_79_scaffold_5.1ta_170wg - Aspergillus nidulans	0	0	22.0	17.2	42.0	18.1	0	01 METABOLISM	
Pc13g13920	hypothetical protein	0	0	25.0	15.4	38.2	12.0	0		
Pc13g14650	strong similarity to hypothetical protein 1166_scaffold_2.1ta_100wg - Fusarium graminearum	2e-11	AE016944_40	product: "phosphoglycolate phosphatase"; Bacteroides thetaiotaomic	126.0	120.0	420.0	13.7	0	
Pc13g15620	strong similarity to hypothetical protein An09g04440 - Aspergillus niger	2e-17	NC081010_14	gene: "BD0A1140"; product: "hypothetical protein"; Neurospora crassa	67.4	36.0	76.6	28.5	0	
Pc13g15630	strong similarity to hypothetical protein An09g04380 - Aspergillus niger	0	0	81.7	20.3	59.3	22.7	0		
Pc13g16650	similarly to hypothetical protein contig1.1ta_1600cg - Aspergillus fumigatus	0	0	441.5	184.3	473.0	186.3	1e-112	99 UNCLASSIFIED PROTEINS	
Pc13g16710	weak similarity to spore-wall fungal hydrophobin deW-A - Aspergillus nidulans	6e-07	S67924	spore-wall fungal hydrophobin deW-A - Emericella nidulans	59.7	13.0	58.6	12.0	1e-87	06.13 proteolytic degradation
Pc14g00580	strong similarity to hypothetical membrane protein YDL237e - yeast [Saccharomyces cerevisiae]	2e-26	S67801	probable membrane protein YDL237e - yeast [Saccharomyces cerevisiae]	248.9	149.9	245.9	111.6	0	
Pc15g00470	weak similarity to hypothetical protein UNC-69 - Caenorhabditis elegans	2e-06	AE030576_35	product: "CG33196-PB"; Drosophila melanogaster chromosome 2L	30.7	12.0	30.1	12.0	0	
Pc15g1120	weak similarity to hypothetical protein SCF34.07 - Streptomyces coelicolor	0.0	AY050511_1	gene: "TmM4"; product: "alpha-L-rhamnosidase A"; Thermomonas	38.2	11.0	63.0	18.4	0	
Pc16g00180	strong similarity to hypothetical protein contig10.1ta_220cg - Aspergillus fumigatus	3e-11	AP005811_119	Gloebacteriaceae PCC 7421 DNA, complete genome, section 1	18.1	12.0	68.0	12.0	0	
Pc16g04550	hypothetical protein	0	0	25.8	12.0	100.3	12.0	0		
Pc16g04840	hypothetical protein	0	0	85.7	12.0	143.1	12.0	0		
Pc16g05400	weak similarity to hypothetical protein contig9.1ta_1510wg - Aspergillus fumigatus	0	0	16.3	12.0	29.6	12.0	0		
Pc16g06660	weak similarity to hypothetical protein encoded by CG4950 - Drosophila melanogaster	0	0	155.8	207.4	469.0	12.0	0		
Pc16g06980	similarly to spore-wall fungal hydrophobin deW-A - Aspergillus nidulans	0	0	70.6	27.0	57.5	13.0	0		
Pc16g08570	strong similarity to hypothetical protein An07g02040 - Aspergillus niger [putative sequencing error]	0	0	27.4	12.0	38.1	12.0	6e-43	01.20 secondary metabolism	
Pc16g08820	similarly to hypothetical protein contig_1_7_scaffold_1.1ta_740cg - Aspergillus nidulans	0	0	105.5	42.1	74.0	53.7	1e-40	99 UNCLASSIFIED PROTEINS	
Pc16g10190	strong similarity to hypothetical glutathione S-transferase SPCC085.07c - Schizosaccharomyces pombe	5e-42	T41222	probable glutathione S-transferase - fission yeast [Schizosaccharom	120.0	120.0	25.4	12.0	0	
Pc16g11150	strong similarity to hypothetical protein contig_1_135_scaffold_11.1ta_750wg - Aspergillus nidulans	9e-09	T41383	hypothetical protein SPCC550.08 - fission yeast [Schizosaccharom	121.4	30.9	92.7	25.9	1e-86	04.05.01.04 transcriptional control
Pc16g11310	strong similarity to allergen Asp I 4 - Aspergillus fumigatus	4e-75	AFRA5PF4_1	gene: "rasp I 4"; product: "Asp I 4"; Aspergillus fumigatus mRNA fo	14.0	12.0	25.5	12.0	0	
Pc16g12230	strong similarity to hypothetical protein contig6_pact.1ta_240wg - Aspergillus fumigatus	1e-151	BRX49607_4	product: "zinc finger protein, putative"; Aspergillus fumigatus BAC p	105.0	39.3	142.2	64.3	0	
Pc16g13260	similarly to hypothetical protein An07g10590 - Aspergillus niger	0	0	67.0	38.3	97.3	12.0	0		
Pc16g13330	similarly to hypothetical protein An05g07000 - Aspergillus niger	0	0	556.9	770.5	1407.2	498.4	1e-86	01.01.01.11 biosynthesis of the cysteine-aromatic group	
Pc16g13500	weak similarity to integral membrane protein PTH11 from patent WO9913094-A2 - Magnaporthe grisea	7e-09	AF119670_1	gene: "PTH11"; product: "integral membrane protein"; Magnaporthe	24.0	28.0	60.0	14.0	0	
Pc16g13220	strong similarity to hypothetical protein contig1490_3.1ta_300cg - Aspergillus fumigatus	8e-51	D09075	sarcosine oxidase (EC 1.5.3.1) - Bacillus sp.	50.8	44.0	76.9	30.3	2e-81	99 UNCLASSIFIED PROTEINS
Pc16g04570	weak similarity to hypothetical protein C0330w - Plasmodium falciparum	2e-13	T19435	hypothetical protein C0330w - malaria parasite Plasmodium falcipa	342.2	58.3	330.0	17.0	0	
Pc16g04650	similarly to hypothetical protein SPAC15A10.00c - Schizosaccharomyces pombe	2e-16	NCB11H24_7	gene: "B11H24.070"; product: "conserved hypothetical protein"; Neu	49.3	17.0	65.0	18.4	0	
Pc20g00300	strong similarity to hypothetical protein An07g02060 - Aspergillus niger	0	0	25.3	15.0	61.4	11.0	0		
Pc20g02790	weak similarity to hypothetical protein contig1487_1.1ta_1190cg - Aspergillus fumigatus	0	0	57.4	25.2	66.9	25.8	5e-54	01.07.10 transport of vitamins, cofactors, and prosthetic groups	
Pc20g03380	hypothetical protein	0	0	1298.5	186.6	2297.3	334.7	1e-45	10 CELLULAR COMMUNICATION/SIGNAL TRANSDUCTION MECHANISM	
Pc20g03740	strong similarity to superoxide dismutase (Cu-Zn) 4-2 - Caenorhabditis elegans	1e-32	JE0098	superoxide dismutase (EC 1.15.1.1) (Cu-Zn) 4-2 - Caenorhabditis el	164.5	16.0	430.7	71.8	0	
Pc20g08040	strong similarity to hypothetical protein contig1487_1.1ta_410wg - Aspergillus fumigatus	5e-31	S69582	hypothetical protein YDR527e - yeast [Saccharomyces cerevisiae]	51.0	46.2	69.2	26.5	0	
Pc20g09550	strong similarity to hypothetical methyltransferase AAO34671.1 - Gibberella zeae	9e-80	AF33639_1	gene: "OHF"; product: "putative methyltransferase"; Fusarium sporo	12.0	12.0	34.1	12.0	0	
Pc20g10840	similarly to hypothetical protein An06g06900 - Aspergillus niger	0	0	52.4	13.7	39.0	16.9	0		
Pc20g11480	hypothetical protein	0	0	581.1	358.3	778.7	315.2	0		
Pc20g11490	hypothetical protein	0	0	250.9	327.8	651.4	119.7	2e-44	01.01.10 amino acid degradation (catabolism)	
Pc20g13710	weak similarity to hypothetical protein - lasmodium falciplanum	4e-08	PFMAL3P7_35	gene: "MAL3P7.34"; "PFC1015c"; product: "hypothetical protein"; F	47.8	12.0	54.5	12.0	0	
Pc20g15760	strong similarity to hypothetical asidase Z37509 - Saccharomyces cerevisiae	1e-106	S50309	hypothetical protein (MEU4.3 region) - yeast [Saccharomyces cerev	16.3	32.4	16.3	2e-78	06.04 protein targeting, sorting and translocation	
Pc21g00950	similarly to hypothetical protein contig_1_168_scaffold_14.1ta_260wg - Aspergillus nidulans	0	0	700.0	148.6	436.1	328.0	2e-69	11.07 detoxification	
Pc21g02340	similarly to hypothetical protein contig127.1ta_340cg - Aspergillus fumigatus	7e-13	AY075446_1	gene: "CG17419"; product: "RE06111p"; Drosophila melanogaster R	12.0	12.0	53.4	12.0	9e-91	04.05.01.04 transcriptional control
Pc21g03010	strong similarity to molasses resistancy protein Rmt1 - Saccharomyces cerevisiae	5e-32	S59140	RTM1 protein - yeast [Saccharomyces cerevisiae] (strain F1100)	104.9	94.0	612.4	96.7	7e-83	99 UNCLASSIFIED PROTEINS
Pc21g06630	similarly to hypothetical protein contig12.1ta_950wg - Aspergillus fumigatus	1e-10	T48778	hypothetical protein 13E11.2600 [imported] - Neurospora crassa	12.0	12.0	27.1	12.0	0	
Pc21g06950	strong similarity to hypothetical protein YML070w - Saccharomyces cerevisiae	5e-41	NC12344_27	gene: "12344.290"; product: "conserved hypothetical protein"; Neu	52.4	13.7	39.0	16.9	0	
Pc21g07580	weak similarity to hypothetical protein An04g09900 - Aspergillus niger	0	0	26.0	12.0	26.7	12.0	0		
Pc21g08140	strong similarity to hypothetical protein contig_1_107_scaffold_7.1ta_1090cg - Aspergillus nidulans	6e-09	AE008918_1	gene: "STM2567"; product: "Gibby-1 prophage protein"; Salmonella	171.8	34.0	229.4	65.3	0	
Pc21g08370	similarly to hypothetical protein An107840 - Aspergillus niger	4e-04	EX089111_15	gene: "HAH7.150"; product: "conserved hypothetical protein"; Neu	19.3	12.0	25.5	12.0	2e-26	01.01.01.01.02 biosynthesis of the glutamate group (proline, hydroxyproline, arginine, glutamine, glutamate)
Pc21g08460	strong similarity to hypothetical protein contig1492_0.1ta_2410wg - Aspergillus fumigatus	0	0	456.3	185.4	595.9	446.9	0		
Pc21g08470	similarly to hypothetical protein 1485_scaffold_9.1ta_210wg - Fusarium graminearum	2e-13	NC081010_20	gene: "BD0A10.200"; product: "conserved hypothetical protein"; Neu	44.3	20.8	41.2	25.8	0	
Pc21g09590	strong similarity to hypothetical protein contig5_pact.1ta_220cg - Aspergillus fumigatus	0	0	116.3	12.0	57.4	28.9	0		
Pc21g09840	similarly to hypothetical protein nc05632.1 - Neurospora crassa	0	0	15.8	12.0	65.6	21.5	0		
Pc21g10690	strong similarity to hypothetical protein contig1488_2.1ta_1690cg - Aspergillus fumigatus	0	0	80.2	38.4	69.3	45.2	1e-154	01.20.37.03 biosynthesis of peptide antibiotics	
Pc21g12310	hypothetical protein	0	0	710.5	518.4	1034.5	328.8	0		
Pc21g12840	strong similarity to hypothetical protein contig_1_153_scaffold_12.1ta_490wg - Aspergillus nidulans	3e-96	MAPEPSYNT_1	gene: "pesA"; product: "peptide synthetase"; Metarhizium anisopliae	258.6	54.7	225.3	94.8	1e-141	14.04.03.05 sporulation and germination
Pc21g14100	strong similarity to hypothetical protein An14g02980 - Aspergillus niger	0	0	36.6	14.5	72.2	12.0	0		
Pc21g15320	similarly to pyoverdine biosynthesis protein PvcA - Pseudomonas aeruginosa	2e-23	H63363	pyoverdine biosynthesis protein PvcA PA2254 [imported] - Pseudon	12.7	12.0	62.4	12.0	0	
Pc21g16130	similarly to hypothetical protein contig1495_1.1ta_1330wg - Aspergillus fumigatus	0	0	171.5	101.3	337.2	36.4	2e-44	03.03.01 mitotic cell cycle and cell cycle control	
Pc21g16440	strong similarity to hypothetical yellowish-green 1 ayg1 - Aspergillus fumigatus	1e-161	AF116902_1	gene: "ayg1"; product: "yellowish-green 1"; Aspergillus fumigatus ye	53.4	12.0	55.5	33.7	0	
Pc21g16070	weak similarity to suppressor of cdc25 mutations Tbt1 - Saccharomyces cerevisiae	3e-06	T16843	TF51 protein - yeast [Saccharomyces cerevisiae]	61.6	57.6	118.2	18.8	7e-53	01.03 nucleotide metabolism
Pc21g01780	strong similarity to hypothetical precursor of spore coat protein spB6 - Neurospora crassa	5e-43	T51044	related to spore coat protein SPB6 precursor [imported] - Neurospor	55.4	12.0	58.7	12.0	0	
Pc22g00460	strong similarity to hypothetical protein An06g1070 - Aspergillus niger	6e-19	BC045017_1	product: "Similar to RIKEN cDNA 3732409C05 gene"; Xenopus lae	62.9	17.2	227.7	87.9	0	
Pc22g02620	strong similarity to hypothetical protein contig12.1ta_1430wg - Aspergillus fumigatus	0	0	545.9	115.5	527.4	445.7	7e-66	11.05.01 resistance proteins	
Pc22g01700	strong similarity to hypothetical protein SCF91.02c - Streptomyces coelicolor	1e-154	SC039306_93	gene: "SC03042"; "SCF91.02a"; product: "conserved hypothetical p	12.5	19.7	27.2	12.0	0	
Pc22g07760	strong similarity to hypothetical protein 1194_scaffold_2.1ta_330wg - Fusarium graminearum	7e-67	NC149_4	gene: "1A9.500"; product: "conserved hypothetical protein"; Neuros	20.9	12.0	40.9	16.0	4e-44	01.05 C-compound and carbohydrate metabolism
Pc22g08820	hypothetical protein	0	0	33.1	12.0	88.4	21.5	0		
Pc22g13780	weak similarity to hypothetical myosin-actin-dehydrogenase spC8 - Streptomyces spectabilis	2e-09	D71201	hypothetical protein PH18B1 - Pyrococcus horikoshii	55.6	18.7	44.1	21.6	0	
Pc22g14300	similarly to hypothetical protein An02g09010 - Aspergillus niger	3e-05	AY184388_1	gene: "MucD"; product: "MUCD"; Mus musculus MUCD (Muc6) gen	132.4	23.1	390.3	15.0	3e-71	06.04 protein targeting, sorting and translocation
Pc22g17040	hypothetical protein	0	0	129.0	17.3	243.9	12.0	3e-29	13.07 cell adhesion	
Pc22g17420	strong similarity to hypothetical protein contig_1_153_scaffold_12.1ta_500cg - Aspergillus nidulans	2e-10	AY159855_1	product: "ankyrin repeat protein EA.2"; Synthetic construct ankyrin	42.0	12.0	68.1	12.0	0	
Pc22g17690	strong similarity to rAsp I 7 - Aspergillus fumigatus	2e-30	AFA3315_1	gene: "rasp I 7"; product: "Asp I 7"; Aspergillus fumigatus mRNA fo	961.9	802.6	1740.9	895.0	0	
Pc22g21690	hypothetical protein	0	0	38.5	12.0	48.2	17.8	1e-145	01.20.05.11 biosynthesis of polyketides	
Pc22g22540	strong similarity to hypothetical protein contig_1_7_scaffold_1.1ta_30cg - Aspergillus nidulans	0	0	46.0	24.0	44.7	12.0	0		
Pc22g22580	strong similarity to hypothetical protein contig_1_153_scaffold_12.1ta_490wg - Aspergillus nidulans	2e-92	AF469045_1	gene: "tex11"; product: "nonribosomal peptide synthetase"; Hypocry	12.0	12.0	33.1	12.0	2e-66	01.05 C-compound and carbohydrate metabolism
Pc22g22620	weak similarity to hypothetical methyltransferase AAO34671.1 - Gibberella zeae	1e-09	AY437642_2	Leptosphaeria maculans ATP binding cassette transporter (ABC4) a	12.0	12.0	50.6	12.0	0	
Pc22g22610	strong similarity to sulphydryl oxidase Sox from patent EP065172-A1 - Aspergillus niger	1e-129	A78767_1	unassigned ORF; Sequence 1 from Patent EP065172.	571.6	311.8	655.2	55.5	0	
Pc22g22920	strong similarity to hypothetical protein contig_1_153_scaffold_12.1ta_50wg - Aspergillus nidulans	0	0	12.0	12.0	33.2	12.0	0		
Pc22g24690	weak similarity to hypothetical protein mlt2143 - Mesorhizobium loti	1e-07	AL646075_14	gene: "RSc3377"; "RS02651"; product: "CONSERVED HYPOTHETI	24.5	44.4	203.4	23.7	0	
Pc22g24750	similarly to hypothetical protein B8J22.03c - Neurospora crassa	6e-18	BRJ2627_2	gene: "B8J22.030"; product: "conserved hypothetical protein"; Neu	12.0	12.0	25.7	12.0	0	
Pc22g24850	similarly to hypothetical protein B8J22.03c - Neurospora crassa	7e-13	BRJ2627_2	gene: "B8J22.030"; product: "conserved hypothetical protein"; Neu	35.5	41.0	217.0	18.3	0	
Pc22g25460	hypothetical protein	0	0	574.7	36.9	1275.2	188.0	0		
Pc22g26260	hypothetical protein	0	0	47.9	16.5	63.8	12.4	0		
Pc22g03740	weak similarity to protein CG33196-PB - Drosophila melanogaster [truncated ORF]	3e-05	AE030576_35	product: "CG33196-PB"; Drosophila melanogaster chromosome 2L	308.4	109.1	644.5	379.8	0	
Pc24g00300	hypothetical protein	0	0	32.6	12.7	43.7	14.5	3e-94	03.03.01.01.09 G2/M transition of mitotic cell cycle	
Pc24g00920	hypothetical protein	0	0	307.9	70.3	231.1	42.8	0		
Pc24g11580	weak similarity to polycystic kidney disease 14-ke 3 - Mus musculus	8e-10	AY164486_1	gene: "Pktd13"; product: "polycystic kidney disease 14-ke 3"; Mus	69.2	14.4	41.8	13.3	0	

Supplementary Table 15. K-mean cluster 7

@=Values given are the average of three independent experiments

orf code	Description of putative <i>P. chrysogenum</i> ORF	Best blast homolog to putative <i>P. chrysogenum</i> ORF			Average transcript levels @				FunCat (auto)	
		p-value	Gene code	description	WIS - PAA	WIS + PAA	DS - PAA	DS + PAA	pval	category
Pc12g020	strong similarity to glucose transporter rco-3 - <i>Neurospora crassa</i>	1e-151	BX842632_4	gene: "B5K2.040"; product: "RCO3-probable glucose transporter";	2281.3	1010.5	113.3	87.6	3e-84	01.05.07 C-compound, carbohydrate transport
Pc12g108	strong similarity to endo 1,5-alpha-arabinanase abnA - <i>Aspergillus niger</i>	7e-82	ABNA_ASPNG	ARABINAN ENDO-1,5-ALPHA-L-ARABINOSIDASE A PRECURSOR (EC 3.2.1.99) (I	1101.3	1179.8	133.8	12e-94	01.05.07 C-compound, carbohydrate transport	
Pc15g000	strong similarity to sugar transporter SUT2 - <i>Pichia stipitis</i>	2e-44	S50708	I hexose transport protein HXT9 - yeast (Saccharomyces cerevisiae)	780.3	199.3	12.0	13.5	0	
Pc15g004	strong similarity to hexose transporter ghp2p - <i>Schizosaccharomyces pombe</i>	0.0	AY081847_1	gene: "mslC"; product: "monosaccharide transporter";	760.9	250.7	189.9	52.1	1e-107	01 METABOLISM
Pc16g131	strong similarity to hypothetical glucan beta-1,3 exoglucanase exgS - <i>Aspergillus phenolicus</i>	0.0	BX849807_83	gene: "exgP"; product: "exo-1,3-beta-D-glucanase, putative";	238.8	107.6	76.0	26.1	0	
Pc16g150	strong similarity to chitinase chnC - <i>Aspergillus nidulans</i>	1e-101	AJ617333_1	I product: "chitin binding protein"; Pichia acaciae plasmid pPact-2 ORF1, ORF2, OR	74.8	55.4	12.0	12.0	1e-102	01.05.01 C-compound and carbohydrate utilization
Pc18g001	weak similarity to isoflavone reductase IFR - <i>Medicago sativa</i>	1e-173	EN134256_1	gene: "cipA"; product: "CipA protein"; Emericella nidulans cipA gene	212.6	187.5	21.9	23.0	0	
Pc18g008	strong similarity to acetate kinase like protein An02g06420 - <i>Aspergillus niger</i>	1e-123	BX842635_4	gene: "B12J7.040"; product: "related to acetate kinase"; Neurospora crassa DNA lin	106.3	61.5	34.5	12.0	0	
Pc18g042	similarity to gentamicin resistance gene like protein An08g01130 - <i>Aspergillus niger</i>	0	0	0	212.7	239.1	75.5	38.1	1e-126	01 METABOLISM
Pc20g028	strong similarity to glutaminase A gtaA - <i>Aspergillus oryzae</i>	0.0	AB029553_1	gene: "gtaA"; product: "glutaminase A"; Emericella nidulans gtaA gene for glutaminas	1177.7	1257.7	246.6	12e-169	13 REGULATION OF / INTERACTION WITH CELLULAR ENVIRONMENT	
Pc21g078	strong similarity to 4-coumarate-CoA ligase 4CL - <i>Arabidopsis thaliana</i>	0.0	AY254391_1	gene: "4CL"; product: "4-coumarate:CoA ligase"; Arabidopsis thaliana 4CL gene, complete	274.7	116.8	23.7	17.9	1e-163	13 REGULATION OF / INTERACTION WITH CELLULAR ENVIRONMENT
Pc21g111	strong similarity to triacylglycerol lipase lipI - <i>Geotrichum candidum</i>	1e-120	BX842680_30	gene: "GOC4.300"; product: "related to cholinesterase precursor"; Neurospora crass	462.5	190.0	258.1	23.2	6e-90	01.05.07 C-compound, carbohydrate transport
Pc21g131	strong similarity to cholinesterase 1 ChE1 - <i>Branchiostoma floridae</i>	8e-48	BC058815_1	Mus musculus hypothetical protein LOC234669, mRNA (cDNA clone	263.0	236.1	96.0	30.1	2e-78	01.05.07 C-compound, carbohydrate transport
Pc21g140	strong similarity to hexose transporter Hx2 - <i>Saccharomyces cerevisiae</i>	1e-170	AY081849_1	gene: "mslE"; product: "monosaccharide transporter";	387.7	127.7	12.0	12.0	6e-82	01.05.04 regulation of C-compound and carbohydrate utilization
Pc21g197	strong similarity to high-affinity glucose transporter HGT1 - <i>Kluyveromyces fragilis</i>	0.0	THA269534_1	gene: "gtt1"; product: "glucose transporter"; Trichoderma harzianum mRNA for glucos	367.8	98.4	12.0	12e-111	01.01 amino acid biosynthesis	
Pc21g215	strong similarity to glucose transporter rco-3 - <i>Neurospora crassa</i>	1e-126	AMM151_1	gene: "AmMst1-1"; product: "AmMst1-1"; A. muscaria mRNA for monosaccharide transp	665.7	168.9	88.1	29.8	4e-79	01.05.07 C-compound, carbohydrate transport
Pc22g009	strong similarity to typhophan synthase Trp5 - <i>Saccharomyces cerevisiae</i>	3e-94	A32959	tyryptophan synthase (EC 4.2.1.20) - <i>Neurospora crassa</i>	239.9	333.9	17.7	14.6	1e-171	01.01 amino acid biosynthesis
Pc22g030	strong similarity to sugar transporter SUT2 - <i>Pichia stipitis</i>	1e-164	NC1911_7	gene: "13E11.140"; product: "probable sugar transporter"; Neurospora crassa DNA	1109.3	428.5	42.1	19.7	4e-78	01.01 amino acid metabolism
Pc22g094	similarity to nonribosomal peptide synthase MxaA - <i>Stigmatella aurantica</i>	9e-31	AY495593_1	gene: "PKS3"; product: "polyketide synthase"; Gibberella moniliformis polyketide syn	696.1	129.9	108.3	21.4	3e-51	01.03.16 polynucleotide degradation
Pc22g122	strong similarity to choline permease Hxm1 - <i>Saccharomyces cerevisiae</i>	2e-51	EN1131668_1	gene: "gabaA"; product: "GABA permease"; Emericella nidulans gabaA gene	139.6	85.9	100.2	12e-3e-51	01.03.16 polynucleotide degradation	
Pc22g000	strong similarity to ribonuclease H1 3-5 exonuclease like protein An1g03420 - <i>Aspergillus niger</i>	4e-17	AY080843_1	gene: "AtSg67240"; product: "unknown protein"; Arabidopsis thaliana unknown protei	364.1	316.2	53.3	51.0	1e-70	04.05.01.04 transcriptional control
Pc22g000	strong similarity to ribonuclease H1 3-5 exonuclease like protein An1g03420 - <i>Aspergillus niger</i>	4e-17	AY080843_1	gene: "AtSg67240"; product: "unknown protein"; Arabidopsis thaliana unknown protei	364.1	316.2	53.3	51.0	1e-158	04 TRANSCRIPTION
Pc22g004	strong similarity to developmental regulatory protein briaA - <i>Aspergillus nidulans</i>	1e-134	AF533070_1	gene: "briaA"; product: "BRLA"; Aspergillus parasiticus BRLA (bria) gene, complete c	299.2	306.6	68.6	22.2	0.0	01.01.01 unspecified signal transduction
Pc22g015	similarity to copper homeostasis protein Cup9 - <i>Saccharomyces cerevisiae</i>	2e-09	AF170065_1	product: "msi2"; Dario neri msi2 mRNA, complete cds.	362.6	439.7	76.3	58.0	0.0	01.01.01 unspecified signal transduction
Pc22g021	strong similarity to retrotransposon Tto1 - <i>Nicotiana tabacum</i>	1e-135	AP002538_10	Oryza sativa (japonica cultivar-group) genomic DNA, chromosome 1, PAC clone:P04	801.6	934.0	267.7	82.9	1e-115	04 TRANSCRIPTION
Pc22g021	strong similarity to retrotransposon Tto1 - <i>Nicotiana tabacum</i>	1e-135	AP002538_10	Oryza sativa (japonica cultivar-group) genomic DNA, chromosome 1, PAC clone:P04	801.6	934.0	267.7	82.9	3e-21	04 TRANSCRIPTION
Pc12g045	weak similarity to Wilms tumor susceptibility protein WT1 - <i>Homo sapiens</i>	9e-50	NCB13020_2	gene: "B13020.050"; product: "related to finger protein AZF1"; Neurospora crassa C	175.3	70.9	53.0	17.4	0	
Pc12g165	weak similarity to bZIP transcription factor Yap5 - <i>Saccharomyces cerevisiae</i>	4e-40	BX842630_4	gene: "B13D15.040"; product: "hypothetical protein"; Neurospora crassa DNA linkag	941.4	823.2	152.1	111.8	3e-26	04 TRANSCRIPTION
Pc13g155	weak similarity to hypothetical transcription factor CAD37059.1 - <i>Neurospora crassa</i>	8e-09	BX849606_08	product: "possible zinc finger protein"; Aspergillus fumigatus BAC pilot project superc	164.3	108.2	40.6	17.8	0.0	10.01.01 unspecified signal transduction
Pc22g041	similarity to 137 kD subunit of DNA-directed RNA polymerase I Rpa12 - <i>Saccharomyces cerevisiae</i>	1e-11	A46107	DNA-directed RNA polymerase (EC 2.7.7.6) I chain A12.2 - yeast (Saccharomyces i	111.3	72.4	32.0	14.6	0	
Pc24g019	strong similarity to retrotransposon Tto1 - <i>Nicotiana tabacum</i>	1e-123	AO179899_11	gene: "OSJNB0092M19.11"; product: "gag-pol polyprotein"; Oryza sativa chromos	391.1	395.6	125.5	47.5	1e-41	03.03.05.03 cell cycle dependent actin filament reorganization
Pc22g000	strong similarity to aspergillopepsin II precursor (acid proteinase A) - <i>Aspergillus niger</i>	7e-64	A41025	aspergillopepsin II (EC 3.4.23.19) precursor - Aspergillus niger (var. macrosporus)	438.8	67.6	114.2	21.2	6e-39	06.04 protein targeting, sorting and translocation
Pc12g075	similarity to coflin Cof1 - <i>Saccharomyces cerevisiae</i>	1e-21	NCB208_27	gene: "B208.270"; product: "related to coflin"; Neurospora crassa DNA linkage grou	474.3	84.7	31.4	12e-86	01.06.03 proteolytic degradation	
Pc16g150	similarity to ankryrin like protein An11g03610 - <i>Aspergillus niger</i>	4e-09	AF017257_17	product: "ankryrin repeat domain protein"; Wolbachia endosymbiont of Drosophila me	82.4	76.0	12.0	12e-54	01.05.07 C-compound, carbohydrate transport	
Pc21g023	strong similarity to aspergillopepsin apnS - <i>Aspergillus phenolicus</i>	8e-93	AF439995_1	product: "pepsin-type protease"; Talaromyces emersoni pepsin-type protease gene,	100.5	56.7	15.0	12.0	0	
Pc21g130	strong similarity to membrane protein Tpa2 - <i>Saccharomyces cerevisiae</i>	0.0	AJ515522_1	Zygosaccharomyces bailii ftz1 gene for fructose facilitator	556.4	103.9	12.0	12.0	0	
Pc21g227	strong similarity to integral membrane protein like protein An04g06980 - <i>Aspergillus niger</i>	8e-08	BX849605_15	product: "integral membrane protein, putative"; Aspergillus fumigatus BAC pilot proje	239.7	51.1	18.4	12.0	2e-54	01.05.04 regulation of C-compound and carbohydrate utilization
Pc20g075	strong similarity to 1,4-beta-D-arabinoxylan arabinofuranohydrolase axhA - <i>Aspergillus niger</i>	1e-133	ANXHA_1	gene: "axhA"; product: "(1,4)-beta-D-arabinoxylan arabinofuranohydrolase"; A niger	179.9	108.3	24.1	12e-106	13.11.03.07 pheromone response	
Pc22g235	strong similarity to cercosporin transporter CFP - <i>Cercospora kikuchii</i>	1e-175	AF282225_1	gene: "Bcmf1"; product: "DHA1A-like major facilitator"; Botryotinia fuckeliana DHA1	874.3	712.6	373.5	42.5	0	
Pc12g164	strong similarity to cell polarity protein tea1p - <i>Schizosaccharomyces pombe</i>	0.0	AJ628227_1	gene: "teaA"; product: "kelch-domain protein"; Emericella nidulans teaA gene for kelc	668.5	482.1	10.8	86.4	0	
Pc21g183	strong similarity to rodless protein rdaA - <i>Aspergillus nidulans</i>	7e-39	AFHYDROF_1	gene: "HYPI"; product: "hydrophobin"; Aspergillus fumigatus hydrophobin (HYP1) ge	436.9	126.4	107.0	37.3	0	
Pc21g237	strong similarity to rodless protein rdaA - <i>Aspergillus nidulans</i>	1e-11	A40323	Rodless protein - Emericella nidulans	1516.4	803.6	620.5	26.9	0	
Pc21g242	strong similarity to rodless protein rdaA - <i>Aspergillus nidulans</i>	4e-50	AFHYDROF_1	gene: "HYPI"; product: "hydrophobin"; Aspergillus fumigatus hydrophobin (HYP1) ge	1500.1	355.1	77.3	14.1	1e-160	10 CELLULAR COMMUNICATION/SIGNAL TRANSDUCTION MECHANISM
Pc16g148	strong similarity to retrotransposon element like protein An08g11510 - <i>Aspergillus niger</i> [putative sequenc	5e-32	A35049	ankryrin 1, erythrocyte splice form 2 - human	157.2	82.0	34.0	19.4	0	
Pc16g094	similarity to ankryrin ANK1 - <i>Homo sapiens</i>	6e-28	NCB7823_13	ankryrin 1, erythrocyte splice form 2 - human	211.4	244.1	12.0	12e-85	01.04.07 phosphate transport	
Pc21g138	strong similarity to transmembrane protein PTH11 - <i>Magnaporthe oryzae</i>	2e-69	S19434	gene: "B7H23.130"; product: "related to L-fucose permease"; Neurospora crassa D	122.5	146.2	53.7	12e-26	01.05.07 C-compound, carbohydrate transport	
Pc06g017	strong similarity to hypothetical transport protein YCR023c - <i>Saccharomyces cerevisiae</i>	1e-18	CNS07TIX_2	probable transport protein YCR023c - yeast (Saccharomyces cerevisiae)	1399.9	1208.9	157.2	146.9	0	
Pc15g017	similarity to hypothetical membrane protein YMR088c - <i>Saccharomyces cerevisiae</i>	0	0	DNA centromeric region sequence from BAC D1P15B03, DP38F06 of chromosome 5	616.6	162.5	248.2	12.0	0	
Pc01g005	weak similarity to hypothetical protein An12g08820 - <i>Aspergillus niger</i>	0	0	0	339.2	344.1	121.5	12.0	0	
Pc04g000	similarity to hypothetical protein contig1492_0.1fa_2370cg - <i>Aspergillus fumigatus</i>	0	0	0	837.0	295.2	276.4	32.4	0	
Pc06g002	similarity to hypothetical protein B208.260 - <i>Neurospora crassa</i>	3e-07	NCB208_14	gene: "B208.140"; product: "conserved hypothetical protein"; Neurospora crassa D	213.9	100.7	12.0	12.0	0	
Pc06g003	strong similarity to hypothetical protein contig1493_1.1fa_1830wg - <i>Aspergillus fumigatus</i>	7e-10	BX321860_127	product: "putative death on curing protein"; Nitrosomonas europaea ATCC 19718, C	457.2	156.1	43.4	12.0	0	
Pc06g003	strong similarity to hypothetical protein contig1493_1.1fa_1830wg - <i>Aspergillus fumigatus</i>	0	0	0	238.6	135.1	65.7	12e-50	01.05 C-compound and carbohydrate metabolism	
Pc06g004	weak similarity to hypothetical protein contig1494_4.1fa_580wg - <i>Aspergillus fumigatus</i>	0	0	0	512.9	761.4	150.2	37.5	0	
Pc06g008	strong similarity to splicing coactivator subunit like protein An01g13270 - <i>Aspergillus niger</i> [truncated ORF]	2e-21	NC6982_1	gene: "69B2.020"; product: "related to glucan 1, 4-alpha-glucosidase"; Neurospora c	294.6	159.6	104.8	42.9	1e-49	04.03.06 RNA modification
Pc06g011	similarity to hypothetical protein 1460_scaffold_8.1fa_310cg - <i>Fusarium graminearum</i>	2e-06	NCB10C3_2	gene: "B10C3.020"; product: "hypothetical protein"; Neurospora crassa DNA linkage	263.1	44.6	43.5	22.1	0	
Pc06g016	strong similarity to hypothetical protein contig_1.1fa_170cg - <i>Aspergillus nidulans</i>	6e-47	S48907	probable purine nucleotide-binding protein YHR16bw - yeast (Saccharomyces cerevi	695.8	516.7	131.0	80.0	1e-138	99 UNCLASSIFIED PROTEINS
Pc06g017	hypothetical protein	0	0	0	108.1	80.9	12.0	12.0	0	
Pc09g000	similarity to hypothetical protein YER080w - <i>Saccharomyces cerevisiae</i>	2e-15	S50583	hypothetical protein YER080w - yeast (Saccharomyces cerevisiae)	157.2	152.7	12.0	12e-26	14.04 cell differentiation	
Pc09g000	hypothetical protein	0	0	0	75.3	117.7	12.0	12.0	0	
Pc09g000	weak similarity to hypothetical protein BAB55333.1 - <i>Homo sapiens</i>	2e-07	BC001705_1	Homo sapiens serine active site containing 1, mRNA (cDNA clone	1197.6	624.6	12.0	12e-26	01.04 phosphate metabolism	
Pc09g001	weak similarity to hypothetical protein - <i>Plasmodium falciparum</i> [truncated ORF]	6e-06	PF4929354_65	gene: "PFE1535w"; product: "hypothetical protein"; Plasmodium falciparum strain 3D	1222.8	1118.4	12.0	12.0	0	
Pc12g004	strong similarity to hypothetical protein An09g02520 - <i>Aspergillus niger</i>	5e-09	AY437641_1	gene: "ST1"; product: "serine/threonine kinase"; Leptosphaeria maculans serine/thr	225.6	73.2	47.6	24.6	4e-78	01.05.01.01.01 sugar, glucoside, polyol and carboxylate catabolism
Pc12g008	similarity to hypothetical protein C03392 - <i>Caulobacter crescentus</i>	2e-54	NCB8P8_4	gene: "B8P8.040"; product: "conserved hypothetical protein"; Neurospora crassa D	1974.2	2466.3	18.1	109.0	0	
Pc12g013	strong similarity to hypothetical protein contig31_part_ii.1fa_1360wg - <i>Aspergillus fumigatus</i>	5e-14	AE016938_2	product: "endo-1,4-beta-xylosylase D precursor"; Bacteroides thetaiotaomicron VPI-54	1049.5	1447.0	279.8	54.5	0	
Pc12g028	hypothetical protein	0	0	0	424.5	119.9	179.4	22.6	0	
Pc12g088	similarity to hypothetical protein SPAC1420.01c - <i>Schizosaccharomyces pombe</i> [putative sequencing error]2e-13	0	T37664	hypothetical protein SPAC1420.01c SPAC56E4.08c - fission yeast (Schizosaccharo	357.3	242.9	92.4	46.5	0	
Pc12g118	strong similarity to hypothetical protein contig1493_1.1fa_1830wg - <i>Aspergillus niger</i>	0	0	0	292.6	108.9	18.1	21e-80	04.05.01.04 transcriptional control	
Pc12g121	strong similarity to hypothetical protein 9G6.360 - <i>Neurospora crassa</i>	3e-21	BX849607_60	product: "hypothetical protein, conserved"; Aspergillus fumigatus BAC pilot project su	129.3	125.8	26.4	12.0	0	
Pc12g121	strong similarity to hypothetical zinc-finger protein flbC - <i>Aspergillus nidulans</i>	5e-97	AF083468_1	gene: "flbC"; product: "putative zinc finger protein"; Emericella nidulans putative zinc	103.4	57.4	24.0	14.2	5e-44	20.25.01.03 hearing
Pc13g010	strong similarity to hypothetical protein An14g02490 - <i>Aspergillus niger</i>	0	0	0	150.5	168.9	41.6	27.2	8e-58	01.01.04 regulation of amino acid metabolism
Pc13g025	similarity to cellulosomal scaffold anchoring protein C - <i>Acetivibrio cellulolyticus</i>	4e-19	AY221113_1	gene: "scaC"; product: "cellulosomal scaffold anchoring protein C"; Acetivibrio cell	236.9	137.6	62.4	15.4	4e-83	10.01.01 unspecified signal transduction
Pc13g031	strong similarity to myb-like DNA binding protein flbD - <i>Aspergillus nidulans</i>	1e-75	EN19882_1	gene: "flbD"; product: "flbD"; Emericella nidulans DNA binding protein flbD (flbD) g	279.3	219.4	68.7	31.6	0	
Pc13g044	similarity to hypothetical MAPK protein kinase-like protein ML14.3 - <i>Arabidopsis thaliana</i>	2e-46	NCB11H24_8	gene: "B11H24.090"; product: "conserved hypothetical protein"; Neurospora crassa	252.4	53.4	68.1	14.9	0	
Pc13g044	similarity to hypothetical protein contig_1.105_scaffold_7.1fa_350cg - <i>Aspergillus nidulans</i>	0	0	0	150.1	117.1	45.4	22.5	0	
Pc13g048	hypothetical protein	0	0	0	228.0	92.8	47.1	28.7	1e-116	01.05 C-compound and carbohydrate metabolism
Pc13g073	strong similarity to hypothetical protein An04g02250 - <i>Aspergillus niger</i>	0	0	0	244.7	291.0	67.5	50.1	0	
Pc13g073	strong similarity to hypothetical protein contig_1.93_scaffold_6.1fa_580cg - <i>Aspergillus nidulans</i>	1e-45	BX294142_8	gene: "pef"; product: "pectate lyase"; Pirella sp. strain 1 complete genome; segmen	72.4	195.1	19.6	22.8	0	
Pc13g0										

Pc16g0711 similarity to hypothetical protein YCR010c - Saccharomyces cerevisiae	3e-17	S31258	probable membrane protein FUNG4 - yeast (Saccharomyces cerevisiae)	97.6	99.9	57.3	12.0		01.04.04 regulation of phosphate utilization	
Pc16g085f similarity to hypothetical protein ncu02885.1 - Neurospora crassa	2e-12	TCU74762_1	gene: "tcrk1"; product: "cdc2-related protein kinase 1"; Trypanosoma cruzi cdc2-rel	315.9	262.8	14.3	14.3	0		0
Pc16g087f strong similarity to Pcl-like cyclin pclA - Aspergillus nidulans	1e-102	ANI272133_1	gene: "pclA"; product: "cyclin"; Aspergillus nidulans pclA gene for cyclin	361.6	210.0	64.2	12.0	0		0
Pc16g098f hypothetical protein				260.4	103.5	87.5	24.4	0		0
Pc16g125f strong similarity to hypothetical protein An13g01540 - Aspergillus niger				95.2	68.3	12.0	12.0	3e-50	01.05 C-compound and carbohydrate metabolism	
Pc16g145f weak similarity to cyanovirin-N CV-N - Nostoc ellipsosporum				134.0	243.9	32.3	12.0	1e-46	01.04 phosphate metabolism	
Pc16g150f strong similarity to hypothetical protein ncu05319.1 - Neurospora crassa	4e-12	T29861	hypothetical protein F10G2.5 - Caenorhabditis elegans	174.2	157.1	12.0	12.0	0		0
Pc17g000f hypothetical protein				133.5	24.0	12.0	12.0	0		0
Pc17g003f hypothetical protein				358.2	413.7	111.0	13.6	0		0
Pc18g015f hypothetical protein				781.8	1173.4	687.3	83.1	0		0
Pc18g017f hypothetical protein				736.8	826.2	104.1	12.0	0		0
Pc18g020f strong similarity to hypothetical protein contig1471.1.f1a.1190cg - Aspergillus fumigatus				372.2	280.9	12.0	12.0	0		0
Pc18g027f strong similarity to hypothetical protein contig1488.2.f1a.1200wg - Aspergillus fumigatus				244.9	133.8	30.8	23.3	0		0
Pc18g030f weak similarity to integral membrane protein PTH11 - Magnaporthe grisea	5e-09	NCB7H23_13	gene: "B7H23.130"; product: "related to L-fucose permease"; Neurospora crassa D	311.1	276.0	165.1	19.2	0		0
Pc18g031f strong similarity to hypothetical protein mg00375.1 - Magnaporthe grisea	8e-82	AY157839_1	gene: "gluB"; product: "beta-1,3-glucanase B"; Lysobacter enzymogenes strain N4-7	1069.6	612.0	259.0	26.6	0		0
Pc18g032f weak similarity to hypothetical protein m2143 - Mesorhizobium loti	2e-06	AP002999_29	gene: "mli2143"; Mesorhizobium loti DNA, complete genome, section 6/21.	491.4	270.4	132.3	29.3	0		0
Pc18g045f weak similarity to hypothetical protein An04g07540 - Aspergillus niger				252.2	147.9	354.9	12.0	0		0
Pc18g052f strong similarity to hypothetical protein 53H1.090 - Neurospora crassa	3e-25	BX842633_9	gene: "53H1.090"; product: "hypothetical protein"; Neurospora crassa DNA linkage c	99.9	64.9	12.7	12.0	0		0
Pc20g016f similarity to hypothetical protein An07g02230 - Aspergillus niger				829.2	617.2	118.2	37.6	0		0
Pc20g020f strong similarity to hypothetical protein contig1487.1.f1a.1460cg - Aspergillus fumigatus				633.5	462.2	225.7	64.0	1e-136	99 UNCLASSIFIED PROTEINS	
Pc20g028f hypothetical protein				93.4	52.8	12.0	12.0	3e-35	99 UNCLASSIFIED PROTEINS	
Pc20g027f strong similarity to hypothetical membrane protein YPR157w - Saccharomyces cerevisiae	1e-54	S61141	probable membrane protein YPR157w - yeast (Saccharomyces cerevisiae)	507.2	255.3	100.5	14.5			0
Pc20g070f similarity to hypothetical Ydr124wp-like protein - Pneumocystis carinii	6e-18	AF309805_4	product: "Ydr124wp-like protein"; Pneumocystis carinii f. sp. carinii glutathione synt	1380.0	1037.4	501.9	180.7	2e-76	30.04 cytoskeleton	
Pc20g070f strong similarity to hypothetical protein contig1487.1.f1a.2180cg - Aspergillus fumigatus				184.4	147.1	63.7	26.6	0		0
Pc20g078f strong similarity to hypothetical protein contig_1_60_scaffold_4.f1a.40wg - Aspergillus nidulans	6e-27	BX846907_5	product: "hypothetical protein"; Aspergillus fumigatus BAC pilot project supercontig;	128.7	96.4	12.0	12.0	2e-65	04.05.01.04 transcriptional control	
Pc20g092f weak similarity to hypothetical integral membrane protein SC10B7.28 - Streptomyces coelicolor	1e-101	BX842629_33	gene: "B20J13.330"; product: "conserved hypothetical protein"; Neurospora crassa i	456.9	283.2	66.9	91.5	0		0
Pc20g128f similarity to hypothetical protein An08g12140 - Aspergillus niger	1e-05	BC064665_1	gene: "zgc:63970"; product: "hypothetical protein MGC63970"; Danio rerio hypothi	226.1	210.1	89.7	12.0	0		0
Pc21g010f similarity to hypothetical protein ncu01891.1 - Neurospora crassa				229.6	127.9	12.0	12.0	0		0
Pc21g010f weak similarity to hypothetical protein An07g0500 - Aspergillus niger				219.8	205.3	56.5	28.5	0		0
Pc21g032f similarity to hypothetical protein contig_1_187_scaffold_29.f1a.20cg - Aspergillus nidulans				362.2	267.2	56.0	25.5	2e-75	01.06 lipid, fatty-acid and isoprenoid metabolism	
Pc21g053f hypothetical protein				1059.8	777.3	886.9	40.4	0		0
Pc21g068f strong similarity to mono- and diacylglycerol lipase precursor - Penicillium camembertii	1e-156	JQ1188	mono- and diacylglycerol lipase (EC 3.1.1.-) precursor - Penicillium camembertii	1937.4	825.4	114.9	475.3	0		0
Pc21g073f strong similarity to hypothetical protein An18g02150 - Aspergillus niger	9e-20	BX908807_3	gene: "B13B7.030"; product: "putative protein"; Neurospora crassa DNA linkage gro	340.8	334.7	71.3	59.6	4e-57	01.05 C-compound and carbohydrate metabolism	
Pc21g086f weak similarity to hypothetical transcription regulator SPCC4F11.01 - Schizosaccharomyces pombe	2e-38	NCB11C21_4	gene: "B11C21.040"; product: "conserved hypothetical protein"; Neurospora crassa	584.0	695.0	546.9	60.1	0		0
Pc21g091f strong similarity to hypothetical protein contig42.f1a.1900cg - Aspergillus fumigatus	8e-14	T39896	probable nucleic acid-binding protein - fission yeast (Schizosaccharomyces pombe)	219.9	151.1	50.5	18.2	5e-28	01.05 C-compound and carbohydrate metabolism	
Pc21g150f hypothetical protein [putative pseudogene]				79.9	53.0	14.2	12.0	0		0
Pc21g186f strong similarity to hypothetical protein B1D4.110 - Neurospora crassa	5e-35	BX908788_9	gene: "B23N11.090"; product: "conserved hypothetical protein"; Neurospora crassa	836.8	1004.2	215.5	135.0	2e-55	01 METABOLISM	
Pc21g188f similarity to hypothetical protein An02g09830 - Aspergillus niger	4e-05	T30886	integumentary mucin B.1 - African clawed frog (fragment)	62.7	87.4	14.4	12.0	1e-77	01.20.17.03 biosynthesis of amines	
Pc21g206f strong similarity to cDNA GS1 - Homo sapiens [putative sequencing error]	6e-46	T40833	haloacid dehalogenase-like hydrolase - fission yeast (Schizosaccharomyces pombe)	589.8	680.2	47.7	54.8	0		0
Pc21g220f strong similarity to hypothetical protein contig_1_153_scaffold_12.f1a.510wg - Aspergillus nidulans	4e-65	AFU293806_1	gene: "tyr1"; product: "tyrosinase"; Aspergillus fumigatus tyr1 gene for tyrosinase, ex	224.0	90.6	79.0	12.0	0		0
Pc21g221f strong similarity to hypothetical protein An16g00290 - Aspergillus niger	2e-12	NCB19A17_10	gene: "B19A17.100"; product: "hypothetical protein"; Neurospora crassa DNA linka	78.4	49.1	12.0	12.0	7e-64	01.04 phosphate metabolism	
Pc21g229f weak similarity to hypothetical protein An01g11580 - Aspergillus niger				126.0	160.8	12.0	12.0	0.0	99 UNCLASSIFIED PROTEINS	
Pc22g019f strong similarity to hypothetical protein An09g02520 - Aspergillus niger	3e-12	AY437641_1	gene: "ST1"; product: "serine/threonine kinase"; Leptosphaeria maculans serine/thre	421.9	202.0	67.3	26.5	0		0
Pc22g063f strong similarity to hypothetical membrane protein YNL279w - Saccharomyces cerevisiae	2e-60	S63253	probable membrane protein YNL279w - yeast (Saccharomyces cerevisiae)	275.9	210.5	12.0	12.0	0		0
Pc22g132f hypothetical protein				85.5	70.9	13.6	16.7	0		0
Pc22g160f strong similarity to hypothetical protein contig5_part_ii.f1a.1430wg - Aspergillus fumigatus				364.4	375.6	256.4	46.9	0		0
Pc22g165f hypothetical protein				182.6	114.7	12.0	12.0	5e-49	25.05.15 myogenesis	
Pc22g178f strong similarity to hypothetical protein contig40.f1a.650cg - Aspergillus fumigatus				1426.5	1249.7	338.0	197.2	0		0
Pc22g185f weak similarity to polycystic kidney disease 1-like 3 - Mus musculus	7e-09	AY164486_1	gene: "Pkd13"; product: "polycystic kidney disease 1-like 3"; Mus musculus polycys	380.5	235.4	171.1	34.9	8e-42	25.05.10 late embryonic development	
Pc22g244f similarity to hypothetical protein An12g08820 - Aspergillus niger				322.0	305.3	128.1	46.7	0		0
Pc22g254f strong similarity to hypothetical protein An01g05100 - Aspergillus niger	1e-04	T09108	RNA binding protein, 24K, chloroplast - spinach (fragment)	119.1	52.9	22.4	12.0	1e-42	99 UNCLASSIFIED PROTEINS	
Pc22g265f hypothetical protein				486.2	176.7	20.1	12.0	0		0
Pc22g271f similarity to hypothetical Ydr124wp-like protein - Pneumocystis carinii	3e-19	AF309805_4	product: "Ydr124wp-like protein"; Pneumocystis carinii f. sp. carinii glutathione synt	326.2	331.0	12.0	12.0	0		0
Pc23g003f strong similarity to hypothetical protein 1103_scaffold_1.f1a.190cg - Fusarium graminearum				449.9	222.6	65.9	27.3	0		0
Pc23g003f hypothetical protein				12.0	12.0	12.0	15.5	0		0
Pc24g006f similarity to hypothetical protein An15g05640 - Aspergillus niger				975.2	403.9	12.0	12.0	0		0
Pc24g007f strong similarity to hypothetical protein An02g07830 - Aspergillus niger				460.0	103.1	14.0	65.9	0		0
Pc24g008f hypothetical protein				720.7	20.9	12.0	12.0	0		0
Pc24g008f strong similarity to hypothetical protein An11g09490 - Aspergillus niger				330.8	12.0	12.0	12.0	0		0
Pc24g008f hypothetical protein				83.0	12.0	12.0	12.0	0		0
Pc24g014f strong similarity to hypothetical protein An11g09490 - Aspergillus niger				239.0	12.0	12.0	12.0	0		0
Pc24g016f hypothetical protein				1761.1	13.0	22.1	12.0	0		0
Pc24g016f hypothetical protein				236.2	14.3	28.4	17.2	0		0
Pc24g025f hypothetical protein				125.7	97.4	36.2	12.0	0		0
Pc24g027f hypothetical protein				982.9	198.9	91.2	28.6	0		0
Pc36g000f weak similarity to hypothetical membrane protein YIL151c - Saccharomyces cerevisiae				750.5	100.7	83.0	63.1	0		0

Supplementary Table 16. K-mean cluster 8

0 = Values given are the average of three independent experiments

orf code	Description of putative <i>P. chrysogenum</i> ORF	Best blast homolog to putative <i>P. chrysogenum</i> ORF			Average transcript levels #					FunCat (auto)
		[e-value]	[Gene code]	[description]	WIS - PAA	WIS + PAA	OS - PAA	OS + PAA	pval	
Pc00g0430	strong similarity to ribonuclease T2 precursor ribN - Aspergillus oryzae	3e-80	JL0205	Bovine ribonuclease T2 (EC 3.1.27.1) precursor - Aspergillus oryzae	200.9	200.9	200.9	200.9	0	01.05 C-compound and carbohydrate metabolism
Pc00g0800	strong similarity to 1,6,6-tetrahydroxyaphthalene reductase like protein An07g01830 - Aspergillus niger	7e-39	BX040445.1	product: "probable short-chain dehydrogenase"; Bordetella pertussis	114.8	100.5	106.2	51.4	3e-39	01.04.07 phosphate transport
Pc00g1310	strong similarity to bromen 2 protein a02 - Aspergillus fumigatus [putative sequencing error]	1e-161	AN005224.1	gene: "b0A"; product: "Nuclease"; Aspergillus nidulans t1A gene	14.2	34.9	12.0	12.0	1e-54	01.05.04 regulation of C-compound and carbohydrate utilization
Pc00g1400	strong similarity to hypothetical amino transferase SPCC10.02 - Schizosaccharomyces pombe	1e-30	T41146	probable amino transferase (EC 2.3.1.1) - fission yeast	71.7	61.1	14.2	14.2	0	01.05 C-compound and carbohydrate metabolism
Pc12g0230	strong similarity to tartrate transport protein tufB - Agrobacterium vitis	2e-93	NC00A10.5	gene: "b0A10.050"; product: "conserved hypothetical protein"	210.7	233.2	119.3	163.5	9e-73	01.05.07 C-compound, carbohydrate transport
Pc02g1730	strong similarity to 3-hydroxy-3-methylglutaryl-CoA synthase Erg13 - Saccharomyces cerevisiae	1e-159	T49719	probable hydroxy-3-methylglutaryl-CoA synthase [imported] - Ni	42.2	20.3	18.8	12.0	0	01.05 C-compound and carbohydrate utilization
Pc12g1610	strong similarity to phosphoenolpyruvate carboxykinase PEPCK - Saccharomyces cerevisiae	8e-74	S30891	probable carboxykinase PEPCK, mitochondrial - yeast (Sacchar	80.3	39.5	25.1	1e-169	0	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc12g0540	strong similarity to citrate synthase citA - Aspergillus niger	0.0	AN024204.1	gene: "citA"; product: "citrate synthetase"; Aspergillus niger c	665.6	586.5	417.2	260.5	0	01.05.01 C-compound and carbohydrate utilization
Pc12g0650	strong similarity to long-chain-fatty-acid-CoA ligase Faa2 - Saccharomyces cerevisiae	0.0	NC01318.11	gene: "B1318.110"; product: "related to long-chain-fatty-acid	366.5	169.8	224.3	236.5	0	
Pc12g0360	strong similarity to phosphoenolpyruvate carboxykinase PEPCK - Kuyseromyces lactis	2e-48	AY048007.1	gene: "b0C"; product: "phosphoenolpyruvate carboxykinase"	460.4	430.1	550.4	210.9	1e-167	13 REGULATION OF / INTERACTION WITH CELLULAR ENVIRONMENT
Pc12g0940	strong similarity to precursor of liver carboxylesterase like protein An12g05840 - Aspergillus niger	0	0	0	578.8	618.8	225.2	184.3	1e-161	01.06.01 lipid, fatty-acid and isoprenoid biosynthesis
Pc12g1340	strong similarity to cephalosporin esterase - Rhodospirillum rubrum	1e-129	BX048234.28	gene: "B16B8.280"; product: "related to triacylglycerol lipase	159.5	214.3	104.2	68.3	0	04.01.04 rRNA processing
Pc12g1520	strong similarity to aureobactin-resistance protein aurA - Aspergillus nidulans	0.0	AF076691.1	gene: "aurA"; product: "aureobactin-resistance protein"; Em	325.9	314.7	143.2	128.7	1e-101	08 CLASSIFICATION NOT YET CLEAR-CUT
Pc12g1520	strong similarity to exosome complex exosome-like Rps6 - Saccharomyces cerevisiae	1e-147	BX000789.27	gene: "B130A.270"; product: "related to nucleolar 100K poly	104.4	86.1	77.5	41.8	4e-70	01.04 phosphate metabolism
Pc13g0240	strong similarity to 2,4-dihydroxyhept-2-ene-1,7-dioic acid aldolase hpcH - Escherichia coli	5e-63	BX049005.29	product: "2,4-dihydroxyhept-2-ene-1,7-dioic acid aldolase"; A	70.1	31.7	48.2	51.8	3e-37	01.03.16.01 RNA degradation
Pc13g0270	similarity to SR protein-specific kinase SPRK2 - Mus musculus	1e-16	M0262456.1	product: "WW domain binding protein 6"; Mus musculus WW	57.8	44.5	28.3	17.5	0	01 METABOLISM
Pc13g0320	weak similarity to thymosin nucleoside Triase tucH - Staphylococcus hyicus	5e-31	T35827	probable nucleoside - fission yeast (Schizosaccharomyces pom	84.9	100.2	43.8	75.0	0	01.05 C-compound and carbohydrate metabolism
Pc13g0320	strong similarity to acetyl-CoA carboxylase SPAC5E04.04c - Schizosaccharomyces pombe	0.0	SE02000	gene: "acetyl-CoA carboxylase (EC 6.4.1.2) - smut fungus (Ustilago	505.1	462.8	379.1	192.8	3e-43	01 METABOLISM
Pc13g0410	strong similarity to NADP-dependent malate dehydrogenase mdh - Homo sapiens	0.0	AF528895.1	gene: "mdhA"; product: "NADP-dependent malic enzyme"; A	97.9	96.2	81.3	33.4	1e-83	01.01.01 amino acid biosynthesis
Pc13g0410	weak similarity to methyl sterol oxidase Erg25 - Saccharomyces cerevisiae	2e-48	AF346734.1	product: "sterol 6-alpha-methyl oxidase"; Arabidopsis thaliana	103.2	122.1	12.0	3e-38	0	01.02 nitrogen and sulfur metabolism
Pc13g0630	strong similarity to salt tolerance protein Met22 - Saccharomyces cerevisiae	8e-50	NC018384.7	product: "Met22A.060"; product: "related to 312); 5-BISPHOS	65.9	45.1	32.8	24.8	0	
Pc13g0670	weak similarity to 3-oxoadipate enol-lactone hydrolase part of the dual specificity protein pcal - Rhodococcus opacus	2e-08	B87548	hypothetical protein CC2411 [imported] - Caulobacter crescer	113.5	254.1	66.8	90.0	0	01.06.01 lipid, fatty-acid and isoprenoid biosynthesis
Pc13g0101	similarity to cutinase precursor cutA - Fusarium solii	9e-26	AF130784.1	gene: "cut1"; product: "cutinase"; Blumeria graminis cutinase	126.5	144.1	13.3	35.2	1e-105	01 METABOLISM
Pc13g0860	strong similarity to lovastatin di-ketide synthase lovF - Aspergillus terreus	0.0	AY46505.1	gene: "PK55"; product: "polyketide synthase"; Gibberella m	61.9	33.2	41.6	4e-20	0	01.01.04 regulation of amino acid metabolism
Pc13g0620	strong similarity to chitinase 1 precursor cit1 - Coccidioides immitis	1e-158	EN0003.1	gene: "chit1"; product: "chitinase"; Aspergillus nidulans chB	1005.2	671.0	512.2	261.0	0	
Pc13g1310	similarity to dehydrogenase synthetase dehydrogenase aBf - Aspergillus nidulans	2e-06	AF148411.2	gene: "gpf"; product: "furfuralase"; Glarea loquaxensis polyket	645.5	634.4	600.1	91.9	2e-95	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc13g1310	strong similarity to hypothetical endoglucanase IV - Trichoderma reesei	5e-99	BX049006.3	product: "endoglucanase, putative"; Aspergillus fumigatus B4	120.8	70.1	40.5	57.4	1e-154	13 REGULATION OF / INTERACTION WITH CELLULAR ENVIRONMENT
Pc13g1310	strong similarity to delta-12 fatty acid desaturase - Mortierella angusta	0.0	AF192749.1	gene: "odaA"; product: "oleate delta-12 desaturase"; Asperg	1303.5	1302.6	628.5	530.2	6e-74	01.05.04 regulation of C-compound and carbohydrate utilization
Pc13g1330	strong similarity to triacylglycerol lipase Lip1 - Candida rugosa	1e-59	S23448	triacylglycerol lipase (EC 3.1.1.1) - yeast (Yarrowia lipolytica)	69.8	120.3	65.7	35.7	0	01 METABOLISM
Pc13g1590	strong similarity to hexose transporter Hxt2 - Saccharomyces cerevisiae	1e-156	SE05951	sugar transport protein STP1 - yeast (Saccharomyces cerevisi	330.1	157.9	177.9	273.1	1e-71	01.01.04 regulation of amino acid metabolism
Pc14g0740	strong similarity to fatty-acyl-CoA synthase beta chain tat1 - Schizosaccharomyces pombe	0.0	S15999	fatty-acyl-CoA synthase (EC 2.3.1.8) beta chain - yeast (Ya	878.2	836.0	618.3	366.5	1e-141	01.03.05.01 DNA repair
Pc14g0920	similarity to chitinase 1 precursor cit1 - Aspergillus nidulans	1e-11	AF129447.1	gene: "chit1"; product: "chitinase"; Aspergillus nidulans chB	73.0	61.1	20.2	12.0	0	01.05.01.01.09 antibiotic ester compounds degradation
Pc14g1820	strong similarity to poly(ethylene glycol) kinase-3-phosphatase PNKP - Homo sapiens	7e-69	T3242	product: phosphatase - fission yeast (Schizosaccharomyces	55.6	60.5	34.4	25.1	2e-47	01 METABOLISM
Pc14g0640	strong similarity to triacylglycerol lipase Lip4 - Candida rugosa	1e-125	JN0552	triacylglycerol lipase (EC 3.1.1.3) precursor - yeast (Candida	40.1	51.2	63.4	18.7	1e-108	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc14g0700	weak similarity to acetyl-hydrolyase chitinase 1 precursor cit1 - Aspergillus nidulans	2e-57	BX049006.34	product: "chitinase 1 precursor"; Aspergillus nidulans chB	37.1	63.4	63.4	37.1	0	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc14g0720	strong similarity to cytochrome P450 protein protein An03g06720 - Aspergillus niger	1e-82	AF155773.16	gene: "FUM15"; product: "Fum15G"; Gibberella moniformis m	28.5	12.0	12.0	3e-54	0	01.05.07 C-compound, carbohydrate transport
Pc16g1730	strong similarity to glycerol-3-phosphate dehydrogenase precursor related protein An08g00210 - Aspergillus niger	0.0	T49652	glycerol-3-phosphate dehydrogenase precursor related protein	694.2	677.7	350.7	477.0	1e-173	01 METABOLISM
Pc16g0280	strong similarity to allantoate permease Dab1 - Saccharomyces cerevisiae	2e-60	T41604	probable membrane transport protein - fission yeast (Schizos	100.0	100.0	100.0	76.9	3e-58	01.05.01.01 C-compound, carbohydrate catabolism
Pc16g0280	strong similarity to phenylalanine ammonia-lyase Plal - Aspergillus nidulans	1e-109	A306660.1	strong ORF: Sequence 12 from Patent WO0004002	42.0	107.0	20.9	16.8	0	
Pc16g1740	weak similarity to isoflavone reductase IFR - Medicago sativa	6e-14	AB045984.1	Verticillium dahliae mRNA from tomato pathotype-specific DN	160.1	134.2	48.8	43.1	7e-44	01.01.01 amino acid biosynthesis
Pc16g1760	weak similarity to hypothetical oxidoreductase P4530 - Schizosaccharomyces pombe	7e-34	NC93.1	gene: "P453.100"; product: "hypothetical protein"; fission ye	40.3	40.3	12.0	12.0	0	06.10 assembly of protein complexes
Pc16g0650	strong similarity to acyl-CoA reductase like protein An14g0470 - Aspergillus niger	2e-22	NC041466.1	gene: "B104.100"; product: "related to CDG diacylglycerol-4	230.6	160.1	257.2	24.5	2e-25	01.05.01.01.09 antibiotic ester compounds degradation
Pc16g0680	weak similarity to 1,3-beta-glucanoyltransferase gel1 - Aspergillus fumigatus	0.0	AF121133.1	gene: "ENGL1"; product: "beta(1-3)endoglucanase"; Asperg	142.3	31.4	64.5	12.0	2e-62	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc16g0880	similarity to salicylate hydroxylase salA - Pseudomonas putilla	1e-36	BX049044.156	product: "putative monooxygenase"; Bordetella bronchiseptic	223.3	68.6	63.4	32.0	0	01.05.01 C-compound and carbohydrate utilization
Pc16g1110	similarity to triacylglycerol lipase Gpl4 - Propionibacterium acnes	2e-54	AF055497.1	product: "putative triacylglycerol lipase"; Streptococcus avermitil	241.9	205.0	241.9	73.8	0	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc16g1180	strong similarity to alpha,alpha-trehalase treH - Aspergillus nidulans	0.0	T18304	acid trehalase homolog - Emmericella nidulans	710.2	259.3	427.9	260.4	1e-91	01.01 amino acid metabolism
Pc16g1730	strong similarity to fatty-acid subterminal hydroxylase P450b9y - Fusarium oxysporum	2e-78	AB076738.1	gene: "CYP50G3A"; product: "fatty acid hydroxylase"; Asperg	717.6	622.4	672.1	288.0	0	01 METABOLISM
Pc16g1170	strong similarity to agmatinase like protein An14g0610 - Aspergillus niger	1e-148	NC01346.2.5	gene: "B134S.050"; product: "conserved hypothetical protein";	69.8	120.3	65.7	35.7	0	01.06 lipid, fatty-acid and isoprenoid biosynthesis
Pc16g1140	strong similarity to mannitol-oligosaccharide 1,2-alpha-mannosidase like protein An01g2500 - Aspergillus niger	0.0	SS8766	mannitol-oligosaccharide 1,2-alpha-mannosidase (EC 3.2.1.	138.9	95.2	81.5	15.4	0	
Pc16g1440	strong similarity to CDG diacylglycerol-4-epimerase P4530y - Schizosaccharomyces pombe	2e-93	NC01014.2	gene: "P453.100"; product: "related to CDG diacylglycerol-4	231.9	222.2	120.7	85.2	1e-111	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc16g1450	strong similarity to release like protein An02g0280 - Aspergillus niger	2e-20	NC007704.46	DNA centromeric region sequence from BAC/CP15B03.03; DP3	34.5	42.2	12.0	12.0	0	01.01.01.01.09 antibiotic ester compounds degradation
Pc16g0240	strong similarity to precursor of linoleate diol synthase - Gaeanomyces graminis	0.0	AY502073.1	gene: "ppaA"; product: "ps1 prodigiosinase"; Emmeric	338.2	315.4	210.4	97.3	5e-64	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc16g0240	strong similarity to beta-galactosidase like protein An02g00610 - Aspergillus niger [putative sequencing error]	0.0	BX049229.35	gene: "B02J13.350"; product: "conserved hypothetical protein	638.2	490.0	173.8	159.9	0	
Pc16g0240	strong similarity to NAD-dependent D-arabinofuranose 4-epimerase DAB1 - Candida tropicalis	2e-54	AF458074.1	product: "D-arabinofuranose 4-epimerase"; Aspergillus nidulans pta	217.8	217.8	217.8	150.2	1e-162	01.07.01 detoxification involving cytochrome P450
Pc16g0830	strong similarity to mutanase muA - Penicillium purpurogenum	1e-152	AF124481.1	gene: "mutA"; product: "mutanase"; Penicillium purpurogenu	310.1	231.7	108.7	90.3	9e-99	01.01.01 amino acid biosynthesis
Pc02g1290	strong similarity to Fe(II)-dependent alpha-ketoglutarate dioxygenase YLL057c - Saccharomyces cerevisiae	6e-69	S50963	hypothetical protein YLL057c - yeast (Saccharomyces cerevi	773.6	760.3	209.9	168.8	9e-99	01 METABOLISM
Pc02g1660	strong similarity to phosphatidylglycerophosphate kinase Pip1 - Saccharomyces cerevisiae	1e-75	S33558	phosphatidylglycerophosphate kinase (EC 2.7.1.1) PIP5K1 - ye	414.0	414.0	414.0	1e-179	0	01.02.01.07 nitric oxide biosynthesis
Pc02g2250	strong similarity to chitinase - Chelonus sp.	1e-122	AY130007.1	gene: "Chit1"; product: "extracellular chitinase"; Blumeria gr	355.5	189.3	223.2	45.8	0	
Pc02g4310	strong similarity to fatty acid omega-hydroxylase CYP50S - Fusarium oxysporum	0.0	AB076738.1	gene: "CYP50G3A"; product: "fatty acid hydroxylase"; Asperg	87.0	42.0	38.7	38.9	1e-180	01.04 phosphate metabolism
Pc02g1780	strong similarity to choline acetyltransferase ChAT - Aspergillus niger	1e-12	G79494	product: "choline acetyltransferase (EC 2.3.1.6) - fission yeast	20.0	114.4	114.4	20.0	0	
Pc02g7880	strong similarity to phosphatidylinositol-4-phosphate 5-kinase - Schizosaccharomyces pombe	1e-149	T38001	product: phosphatidylinositol-4-phosphate 5-kinase - fission y	97.8	74.8	48.5	36.2	1e-87	01.05.07 C-compound, carbohydrate transport
Pc02g1030	weak similarity to penicillamine synthase - Streptomyces sp.	0	0	0	45.6	21.1	20.3	36.2	2e-83	08 CLASSIFICATION NOT YET CLEAR-CUT
Pc02g1020	strong similarity to glucose transporter isoC - Neurospora crassa	1e-143	BX042632.4	gene: "B02C.040"; product: "RCCO3-probable glucose transpo	53.4	42.8	12.0	12.0	0	01.03 nucleotide metabolism
Pc02g1210	similarity to glucanase transport-inducing protein gfpA - Schizosaccharomyces pombe	4e-25	S50452	hypothetical protein YEL007w - yeast (Saccharomyces cerevi	218.7	236.0	126.1	91.6	0	01.05.01 C-compound and carbohydrate utilization
Pc02g1280	strong similarity to protein involved in glycosylphosphatidylinositol biosynthesis Gp13 - Saccharomyces cerevisiae	1e-151	S64762	probable membrane protein YEL031c - yeast (Saccharomyces	62.8	58.7	36.3	26.8	1e-40	01 METABOLISM
Pc02g1340	strong similarity to isomyl alcohol oxidase hsd - Aspergillus oryzae	1e-177	NC0484.4	gene: "B04C.040"; product: "probable isomyl alcohol oxidase	98.1	88.1	31.5	16.6	0	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc02g1340	weak similarity to methyl sterol oxidase Erg25 - Saccharomyces cerevisiae	3e-08	AE016793.188	product: "conserved hypothetical protein"; Pseudomonas puti	106.7	123.0	12.0	12.0	0	01 METABOLISM
Pc12g1050	strong similarity to NAD-dependent D-arabinofuranose 4-epimerase DAB1 - Candida tropicalis	1e-153	AY387591.1	gene: "arbd"; product: "D-arabinofuranose dehydrogenase"; Emneri	727.7	773.3	355.3	162.7	0	01.20.37 biosynthesis of peptide antibiotics
Pc12g1080	strong similarity to 3-methylcrotonyl-CoA carboxylase (MCC) biotin-containing alpha subunit MCCA - Homo sapiens	0.0	AY387592.1	gene: "mccA"; product: "3-methylcrotonyl-CoA carboxylase b	258.7	257.6	110.1	188.4	4e-57	01.04.07 phosphate transport
Pc12g1710	strong similarity to peptide synthase like protein An04g0280 - Aspergillus niger	0.0	AF460445.1	gene: "tst1"; product: "Trombostamol peptide synthetase"; H	47.8	27.7	45.0	12.0	0	
Pc12g0510	strong similarity to high-affinity nicotinic acid permease Tnat1 - Saccharomyces cerevisiae	1e-128	BX080809.43	gene: "TNE8.430"; product: "related to nicotinamide mononucl	267.2	248.5	119.1	176.3	1e-102	13.11.01 photoreception and response
Pc12g0550	weak similarity to nitrate reductase (NADPH) nadA - Aspergillus niger	0	0	0	21.5	47.5	32.5	12.0	3e-84	02.11 electron transport and membrane-associated energy conservation
Pc12g0550	strong similarity to hypothetical phosphatidyl-acid-esterifying phosphatase A1 B14D6.220 - Neurospora crassa	0	T49470	phosphatidic acid-esterifying phosphatase A1 related protein	159.3	118.4	114.2	47.5	0	01.01.03 polyketide degradation
Pc12g0180	strong similarity to threonine - Galius galus	1e-122	NC17E5.25	gene: "TRES.270"; product: "conserved hypothetical protein";	298.3	407.5	174.4	202.6	0	01.06.01 lipid, fatty-acid and isoprenoid biosynthesis
Pc12g0330	strong similarity to mRNA turnover 4 protein Mtn4 - Saccharomyces cerevisiae	2e-58	T39330	probable 60S acidic ribosomal protein - fission yeast (Schizo	230.2	216.1	177.5	104.1	1e-94	01.05.01 C-compound and carbohydrate utilization
Pc12g0070	strong similarity to polyketide synthase like protein An02g0180 - Aspergillus niger	0.0	AY465022.1	gene: "PK517"; product: "polyketide synthase"; Botrytis cin	71.6	80.0	65.5	25.1	1e-103	01.04.04 regulation of phosphate utilization
Pc12g0250	strong similarity to alpha-1,6-mannosyltransferase Hct1 - Saccharomyces cerevisiae	3e-27	SPAC1006.5	gene: "hct1"; "SPAC1006.05"; S. pombe chromosome 1 (imp	95.9	29.4	36.7	56.3	1e-75	03.03.01.03 cell cycle checkpoints (checkpoints of morphogenesis, DNA-damage, replication, mitotic phase and spindle)
Pc12g0470	strong similarity to cyclin-dependent protein kinase Pkb85 - Saccharomyces cerevisiae	0.0	T49815	product: multifunctional cyclin-dependent kinase						

Pc22g18690	weak similarity to alpha-1,6-mannosyltransferase Hct1 - Saccharomyces cerevisiae [truncated ORF]	4e-19	AF373493.1	product: "putative mannosyltransferase", <i>Parascoccidioides</i>	61.4	35.3	18.3	22.7	7e-92	01.06 C-compound and carbohydrate metabolism
Pc22g24680	strong similarity to high-affinity glucose transporter HGT1 - <i>Kluyveromyces fragilis</i>	1e-172	AH44343.1	gene: "hgt1", product: "glucose transporter", <i>Hypocrea jecori</i>	48.7	57.9	18.5	72.9	4e-84	01.01.10 amino acid degradation (catabolism)
Pc22g2480	strong similarity to pectate lyase pLYA - <i>Aspergillus niger</i>	1e-119	ANZ67531.1	gene: "pLYA", product: "pectate lyase A", <i>Aspergillus niger</i> str.	29.8	31.8	12.0	12.0	0.0	01.06 C-compound and carbohydrate metabolism
Pc22g24900	similarity to salicylate 1-monooxygenase salI - <i>Pseudomonas putilla</i>	8e-19	AD2770	salicylate hydroxylase [imported] - <i>Agrobacterium tumefaciens</i>	50.1	101.6	18.8	30.1	2e-67	01.03 nucleotide metabolism
Pc22g25050	strong similarity to myo-inositol-1-phosphate synthase like protein Ant1g05030 - <i>Aspergillus niger</i>	0.0	NC01094.10	gene: "INO1H1102", product: "probable myo-inositol-1-phosphatase", <i>Aspergillus niger</i>	90.1	619.8	616.3	469.4	0.0	0.0
Pc22g25650	strong similarity to dUTP pyrophosphatase DUT1 - <i>Candida albicans</i>	2e-49	BX0242627.10	gene: "BLL22", product: "probable dUTP pyrophosphatase", <i>Candida albicans</i>	48.1	415.7	26.1	163.3	0.0	0.0
Pc24g2480	similarity to 35.6K acid phosphatase - <i>Aspergillus nidulans</i>	7e-21	PC08104.1	gene: "phoG", product: "PHOG", <i>Penicillium chrysogenum</i> P	37.3	36.2	12.0	12.0	1e-66	01.07.01 biosynthesis of vitamins, cofactors, and prosthetic groups
Pc24g2480	similarity to 35.6K acid phosphatase - <i>Aspergillus nidulans</i>	7e-21	PC08104.1	gene: "phoG", product: "PHOG", <i>Penicillium chrysogenum</i> P	27.3	46.2	12.0	12.0	1e-67	04. TRANSCRIPTION
Pc24g1450	strong similarity to assembly factor of cytochrome c oxidase Cox11 - <i>Saccharomyces cerevisiae</i>	1e-42	NC013818.14	gene: "B13H1140", product: "probable cytochrome c oxidase", <i>Aspergillus niger</i>	93.1	96.1	26.3	32.2	4e-29	01.03.01 purine nucleotide metabolism
Pc21g17100	strong similarity to zinc-finger transcription factor amA - <i>Aspergillus nidulans</i>	1e-123	B61908	RNA-binding protein amA - <i>Emicorticaria nidulans</i>	160.1	84.1	69.3	73.1	1e-166	05.01 ribosome biogenesis
Pc21g1300	strong similarity to mitochondrial ATP synthase subunit 9 tlc31 - <i>Aspergillus nidulans</i>	8e-31	LW1NCA	H+-transferring two-sector ATPase (EC 3.6.3.14) lipid-binding probable thiosomal protein 54 precursor, mitochondrial - yeast	153.4	163.1	726.6	964.6	2e-55	01.21 electron transport and membrane-associated energy conservation
Pc21g1160	strong similarity to mitochondrial ribosomal protein Nam1 - <i>Saccharomyces cerevisiae</i>	4e-46	S50146	gene: "turd", product: "extragenic suppressor of the bimD6 gene", <i>Aspergillus niger</i>	130.2	120.2	88.6	46.8	0.0	01.04 phosphate metabolism
Pc40g00300	H+-transferring two-sector ATPase protein 6 - <i>Penicillium chrysogenum</i>	2e-99	S42271	H+-transferring two-sector ATPase (EC 3.6.3.14) protein 6 - 1	36.0	33.4	12.5	12.0	0.0	01.08.10 regulation of lipid, fatty-acid and isoprenoid metabolism
Pc21g1650	strong similarity to ras pathway interacting protein mole1p - <i>Schizosaccharomyces pombe</i>	1e-151	BNK42632.7	gene: "SPAC637.07", product: "ras pathway interacting protein 1", <i>Schizosaccharomyces pombe</i>	806.8	199.1	527.7	238.7	7e-70	01.04 phosphate metabolism
Pc12g1210	strong similarity to regulator protein Sin3 - <i>Saccharomyces cerevisiae</i>	0.0	BNK42632.7	gene: "B9C2", product: "related to transcription regulation", <i>Schizosaccharomyces pombe</i>	129.5	184.1	75.4	62.1	1e-74	01.04 phosphate metabolism
Pc12g7790	weak similarity to hypothetical serine threonine phosphatase topA - <i>Fenestudactinium istandicum</i>	3e-11	S50713	gene: "B1D14.120", product: "probable dis1-suppressing protein", <i>Aspergillus niger</i>	199.3	196.4	91.2	103.9	1e-157	01.04 phosphate metabolism
Pc12g8830	strong similarity to nucleolin protein like Ant1g05190 - <i>Aspergillus niger</i>	2e-14	T38810	hypothetical protein SPAC4F10.06 - fission yeast (<i>Schizosaccharomyces pombe</i>)	51.5	44.4	36.6	17.7	2e-94	01.04 phosphate metabolism
Pc12g13830	strong similarity to extragenic suppressor of the bimD6 mutation xtdD - <i>Aspergillus nidulans</i>	0.0	AF013580.1	gene: "turd", product: "extragenic suppressor of the bimD6 gene", <i>Aspergillus niger</i>	130.2	120.2	88.6	46.8	0.0	01.04 phosphate metabolism
Pc12g12820	similarity to dis1-suppressing protein kinase dsk1p - <i>Schizosaccharomyces pombe</i>	8e-18	NC01D14.12	gene: "B1D14.120", product: "probable dis1-suppressing protein", <i>Aspergillus niger</i>	35.2	13.7	13.1	14.9	1e-175	04.05.01.04 transcriptional control
Pc12g23240	strong similarity to protein kinase of the MAP kinase kinase 2 Mkk2 - <i>Saccharomyces cerevisiae</i> [truncated ORF][putative sequencing error]	1e-166	AY42722.3	Podoparas aeneus chromosome 7 MAP kinase kinase 1 (M)	487.1	379.2	181.0	188.2	1e-53	01.06 C-compound and carbohydrate metabolism
Pc12g3850	similarity to protein kinase of the MAP kinase kinase 2 Mkk2 - <i>Saccharomyces cerevisiae</i> [truncated ORF][putative sequencing error]	1e-166	AY42722.3	Podoparas aeneus chromosome 7 MAP kinase kinase 1 (M)	487.1	379.2	181.0	188.2	1e-53	01.06 C-compound and carbohydrate metabolism
Pc13g10160	weak similarity to replication protein cdc23p - <i>Schizosaccharomyces pombe</i>	5e-68	BN897676.13	gene: "B8B24.007", product: "related to transcription regulation", <i>Schizosaccharomyces pombe</i>	45.5	48.2	25.3	22.4	0.0	03.01.01 cellular DNA uptake
Pc13g10330	strong similarity to meiosis control kinase rmlp1 - <i>Schizosaccharomyces pombe</i>	2e-94	FS53329.1	product: "Ran1-like protein kinase", <i>Fusarium solani</i> Rant1-lik	446.7	330.6	220.5	121.9	1e-131	03.01.05 DNA recombination and DNA repair
Pc13g11700	strong similarity to cell division control protein cdc14p - <i>Schizosaccharomyces pombe</i>	6e-96	BNK49068.39	gene: "cdc14", product: "cdc complex protein, putative", <i>Aspergillus niger</i>	254.5	238.5	18.2	97.9	1e-101	01.04 phosphate metabolism
Pc13g12210	similarity to cell division control protein cdc14p - <i>Schizosaccharomyces pombe</i>	6e-96	BNK49068.39	gene: "cdc14", product: "cdc complex protein, putative", <i>Aspergillus niger</i>	194.0	151.6	102.4	66.6	1e-74	01.04 phosphate metabolism
Pc13g13180	strong similarity to protein Rad57 - <i>Saccharomyces cerevisiae</i>	1e-121	BNK49065.128	product: "rad57 protein kinase", <i>Aspergillus fumigatus</i> BAC	71.4	68.8	41.8	30.0	0.0	01.05.04 regulation of C-compound and carbohydrate utilization
Pc13g16710	similarity to dis1-suppressing protein kinase dsk1p - <i>Schizosaccharomyces pombe</i>	2e-10	SPBC520.14	gene: "ada3", product: "SPOC3B30.14C", <i>S. pombe</i> chromosome I cos	65.4	69.4	40.4	18.9	0.0	03.03 cell cycle
Pc13g09160	strong similarity to ADA and SAGA histone acetyltransferase subunit Ada3 - <i>Saccharomyces cerevisiae</i>	1e-32	ADA3ADA3A.1	gene: "ada3", product: "histone deacetylase in activation 3 p", <i>Aspergillus niger</i>	31.6	14.9	12.0	12.0	4e-61	01.05.04 regulation of C-compound and carbohydrate utilization
Pc13g14970	strong similarity to hypothetical tubulin alpha-2 chain tubA2 - <i>Aspergillus nidulans</i>	0.0	S13337	tubulin alpha-2 chain - <i>Emicorticaria nidulans</i>	28.0	18.4	12.0	12.0	4e-42	01.04 phosphate metabolism
Pc13g10660	weak similarity to suppressor of tort1 protein Mpt4 - <i>Saccharomyces cerevisiae</i>	3e-39	S48511	MPT4 protein - yeast (<i>Saccharomyces cerevisiae</i>)	780.9	248.7	248.7	248.7	1e-161	03.01.05 DNA recombination and DNA repair
Pc13g11560	strong similarity to guanine nucleotide-releasing factor Uel1 - <i>Saccharomyces cerevisiae</i>	1e-135	AY365418.10	gene: "ada3", product: "histone deacetylase in activation 3 p", <i>Aspergillus niger</i>	306.9	307.9	166.1	204.9	0.0	03.01.05 DNA recombination and DNA repair
Pc13g01820	similarity to ATP-dependent DNA helicase II 8K kD subunit G22P2 - <i>Homo sapiens</i>	2e-84	SPBC543.3	gene: "SPBC543.03", <i>S. pombe</i> chromosome I cosmid c54	385.7	297.3	103.3	80.4	0.0	01.04 phosphate metabolism
Pc23g0750	strong similarity to centromere microtubule-binding protein Cdb5 - <i>Saccharomyces cerevisiae</i>	0.0	CBF5_ASP1U	CENTROMERE/MICROTUBULE BINDING PROTEIN CBF5 (C)	679.1	686.1	436.2	539.6	1e-66	04.05.05.01 regulation of splicing
Pc23g05050	strong similarity to small protein tyrosine phosphatase tip1p - <i>Schizosaccharomyces pombe</i>	1e-31	A33146	gene: "tip1", product: "protein tyrosine phosphatase", <i>Schizosaccharomyces pombe</i>	177.1	177.1	12.0	12.0	1e-67	01.01.04 phosphate metabolism
Pc23g06960	strong similarity to fuzzy-related protein Fzr1 - <i>Mus musculus</i>	0.0	BNX42618.15	gene: "B13M13.150", product: "fuzzy-related protein Fzr1", <i>Aspergillus niger</i>	76.8	70.3	35.1	34.7	2e-81	01.04.03 DNA synthesis and replication
Pc23g0420	strong similarity to protein Ract1 - <i>Saccharomyces cerevisiae</i>	1e-140	CN059545.87	DNA centromeric region sequence from BAC CP2B206, DPS	106.7	107.1	60.0	49.2	5e-50	01.04 phosphate metabolism
Pc23g0490	weak similarity to DNA polymerase kappa TRF4 - <i>Homo sapiens</i>	1e-15	CC2_EMEN	gene: "tip1", product: "protein tyrosine phosphatase", <i>Schizosaccharomyces pombe</i>	189.5	189.5	12.0	12.0	1e-67	01.01.04 phosphate metabolism
Pc23g10270	strong similarity to protein kinase functional homolog of cdc2 nim1 - <i>Aspergillus nidulans</i>	1e-145	CC2_EMEN	CELL DIVISION CONTROL PROTEIN 2 (EC 2.7.1.1) (CYCLIN	27.2	28.8	23.8	12.0	7e-38	03.01.09.05 DNA conformation modification (e.g. chromatin)
Pc23g1460	strong similarity to replication factor C36 kD subunit RFFC3 - <i>Homo sapiens</i>	1e-160	NC18406.46	gene: "B14D6.460", product: "probable replication factor C38	135.5	134.1	85.6	72.2	1e-136	03.03.01 mitotic cell cycle and cell cycle control
Pc23g11490	strong similarity to suppressor of the xtdD1 mutation xtdD1 - <i>Aspergillus nidulans</i>	2e-27	T45518	gene: "B14D6.460", product: "probable replication factor C38	127.9	127.9	12.0	12.0	1e-67	01.01.04 phosphate metabolism
Pc23g19180	strong similarity to cohesin subunit XSA1 - <i>Xenopus laevis</i>	1e-167	NC01H10.5	gene: "3H103.500", product: "related to nuclear protein SA-1", <i>Aspergillus niger</i>	73.3	59.8	36.4	34.6	5e-68	03.01.03 DNA synthesis and replication
Pc21g02020	similarity to lethal2Dentlesslike (tld2) - <i>Drosophila melanogaster</i>	1e-111	T48835	lethal2Dentlesslike related protein [imported] - <i>Neurospora crassa</i>	96.5	90.8	47.4	40.1	1e-100	03.01.03 DNA synthesis and replication
Pc12g14330	strong similarity to DNA polymerase kappa TRF4 - <i>Homo sapiens</i>	1e-45	A40728	gene: "tip1", product: "protein tyrosine phosphatase", <i>Schizosaccharomyces pombe</i>	189.5	189.5	12.0	12.0	1e-67	01.01.04 phosphate metabolism
Pc12g14620	strong similarity to DNA polymerase kappa TRF4 - <i>Homo sapiens</i>	2e-42	SE2544	hypothetical protein SPAC12G12.13c - fission yeast (<i>Schizosaccharomyces pombe</i>)	49.3	35.8	32.3	16.6	0.0	03.01.03 DNA synthesis and replication
Pc12g17340	strong similarity to sp1-GTPase binding protein atg1p - <i>Saccharomyces cerevisiae</i>	1e-49	YB098087.2	gene: "B13B7.020", product: "probable sp1-GTP-binding protein", <i>Aspergillus niger</i>	113.1	89.8	67.0	36.2	4e-67	03.01.03 DNA synthesis and replication
Pc12g11310	strong similarity to RNA polymerase II transcription factor TFIIB - <i>Aspergillus nidulans</i>	0.0	NC013833.1	gene: "B13B7.020", product: "probable sp1-GTP-binding protein", <i>Aspergillus niger</i>	149.8	149.8	12.0	12.0	1e-67	01.01.04 phosphate metabolism
Pc12g14620	strong similarity to transcription factor C subunit ARFC3 - <i>Ankara adenovirus</i>	1e-149	NCB13833.18	gene: "B13B3.180", product: "related to replication factor C d", <i>Aspergillus niger</i>	152.5	145.8	80.9	67.2	7e-93	01.01.04 phosphate metabolism
Pc12g11660	strong similarity to cell division protein rbd4p - <i>Saccharomyces cerevisiae</i>	0.0	NC12344.8	gene: "12344.060", product: "probable cell division control pr", <i>Aspergillus niger</i>	222.7	181.0	97.7	100.6	3e-92	01.06 C-compound and carbohydrate metabolism
Pc23g0780	strong similarity to liver cancer protein 1 - <i>Aspergillus nidulans</i>	1e-16	CE32059.1	gene: "unc-80", product: "UNC-80", <i>Caenorhabditis elegans</i> C. elegans	198.1	198.1	12.0	12.0	1e-67	03.01.04 phosphate metabolism
Pc23g01220	strong similarity to G T mismatch-specific thymine-DNA glycosylase TGO - <i>Homo sapiens</i>	1e-214	7e-214	hypothetical protein SPCC065.05c - fission yeast (<i>Schizosaccharomyces pombe</i>)	77.1	71.1	40.4	32.4	4e-98	03.03.01 mitotic cell cycle and cell cycle control
Pc23g02950	strong similarity to twa2 - <i>Saccharomyces cerevisiae</i>	0.0	SE7208	hypothetical protein YOR304a - yeast (<i>Saccharomyces cerevisiae</i>)	279.1	236.3	150.6	109.8	1e-237	06.07.03 modification by phosphorylation, dephosphorylation
Pc23g0330	strong similarity to cyclin B nine - <i>Aspergillus nidulans</i>	0.0	S22896	cyclin B - <i>Emicorticaria nidulans</i>	28.0	28.0	12.0	12.0	1e-67	01.03.16.01 RNA degradation
Pc23g0330	weak similarity to cell division control protein Cdc31 - <i>Saccharomyces cerevisiae</i>	1e-31	S47549	cell division control protein CDC31 - yeast (<i>Saccharomyces cerevisiae</i>)	187.2	186.2	126.3	79.6	1e-66	04. TRANSCRIPTION
Pc23g0880	strong similarity to RNA binding protein rntDp - <i>Schizosaccharomyces pombe</i>	1e-128	SE5196	probable membrane protein YPL184c - yeast (<i>Saccharomyces cerevisiae</i>)	225.0	180.6	91.0	46.8	1e-107	04. TRANSCRIPTION
Pc23g0870	weak similarity to DNA polymerase V Pds1 - <i>Saccharomyces cerevisiae</i> [truncated ORF]	4e-55	BNK49231.3	gene: "B1014.030", product: "related to DNA polymerase V", <i>Aspergillus niger</i>	150.5	133.9	102.4	54.4	7e-73	03.01.05 DNA recombination and DNA repair
Pc23g0880	strong similarity to DNA polymerase V Pds1 - <i>Saccharomyces cerevisiae</i> [truncated ORF]	4e-55	BNK49231.3	gene: "B1014.030", product: "related to DNA polymerase V", <i>Aspergillus niger</i>	150.5	133.9	102.4	54.4	7e-73	03.01.05 DNA recombination and DNA repair
Pc23g18050	strong similarity to Rad52 homologue MUS11 - <i>Neurospora crassa</i>	1e-123	AY032591.1	gene: "radc", product: "RADCO", <i>Schizosaccharomces RADCO</i>	36.8	27.4	12.0	12.0	0.0	03.03.01 DNA synthesis and replication
Pc23g18050	strong similarity to Dom34-interacting protein like protein Ant1g01900 - <i>Aspergillus niger</i>	0.0	NC01H112.16	gene: "B11N2.160", product: "conserved hypothetical protein", <i>Aspergillus niger</i>	61.4	26.1	64.9	26.2	9e-29	03.03.01 DNA synthesis and replication
Pc23g10110	strong similarity to polymerase I Pti1 - <i>Saccharomyces cerevisiae</i>	1e-103	DUP7A	DNA-directed RNA polymerase (EC 2.7.7.6) alpha - fission yeast (<i>Schizosaccharomyces pombe</i>)	120.9	120.9	12.0	12.0	1e-67	01.01.04 phosphate metabolism
Pc23g04040	similarity to protein required for accurate chromosome transmission Chit2 - <i>Saccharomyces cerevisiae</i>	5e-08	S50340	CHL2 protein - yeast (<i>Saccharomyces cerevisiae</i>)	40.4	37.1	12.9	12.0	1e-109	01.01.04 regulation of amino acid metabolism
Pc23g04040	strong similarity to developmental regulatory protein bIRA - <i>Saccharomyces cerevisiae</i>	4e-37	A26913	regulatory protein bIRA - <i>Emicorticaria nidulans</i>	12.0	12.0	12.0	12.0	3e-38	04.05.05 mRNA processing (splicing, 5', 3'-end processing)
Pc23g12710	weak similarity to neuron esmatin-like regulatory protein tld4 - <i>Neurospora crassa</i>	1e-137	TS1029	gene: "B0A11.020", product: "probable transcription factor", <i>Aspergillus niger</i>	75.1	75.1	12.0	12.0	1e-67	01.01.04 phosphate metabolism
Pc23g13020	strong similarity to SMAD interacting protein like protein Ant1g07230 - <i>Aspergillus niger</i>	2e-52	AF236664.1	gene: "h2IA", product: "hypothetical zinc-finger protein", <i>Aspergillus niger</i>	103.6	102.4	50.6	44.1	3e-68	04.05.01.04 transcriptional control
Pc23g17010	similarity to transcription activator P34 - <i>Saccharomyces cerevisiae</i>	4e-10	A33970	transcription activator P34 - <i>Saccharomyces cerevisiae</i>	90.6	79.1	27.2	35.0	0.0	01.04 phosphate metabolism
Pc12g11050	strong similarity to hypothetical zinc-finger protein tld4 - <i>Neurospora crassa</i>	1e-49	NC01D12.20	gene: "B0D12.200", product: "conserved hypothetical protein", <i>Aspergillus niger</i>	189.6	189.6	12.0	12.0	1e-67	03.03.01 mitotic cell cycle and cell cycle control
Pc12g11400	strong similarity to subunit of transcription initiation factor TRF1 like protein Ant1g05030 - <i>Aspergillus niger</i>	1e-149	SE2469	probable transcription initiation factor tld4 subunit - fission yeast (<i>Schizosaccharomyces pombe</i>)	333.3	272.3	173.8	107.6	3e-96	04. TRANSCRIPTION
Pc12g1400	similarity to zinc-finger protein CZF1 - <i>Candida albicans</i>	5e-09	AY109221.1	gene: "MPP1", product: "2Hf2925 transcription factor", <i>Pic</i>	52.8	48.8	24.8	34.7	3e-37	01.05.04 regulation of C-compound and carbohydrate utilization
Pc12g14680	similarity to DNA binding regulatory protein RGA1 - <i>Aspergillus nidulans</i>	1e-11	SC0264.1	gene: "B0B11.020", product: "probable zinc-finger protein", <i>Aspergillus niger</i>	129.1	129.1	12.0	12.0	1e-67	01.01.04 phosphate metabolism
Pc12g17670	similarity to nuclear polyadenylated-RNA-binding protein Nab3 - <i>Saccharomyces cerevisiae</i>	3e-18	T38767	probable RNA-binding protein - fission yeast (<i>Schizosaccharomyces pombe</i>)	83.1	57.4	49.9	24.3	2e-31	01.01.04 regulation of amino acid metabolism
Pc12g1650	strong similarity to zinc-finger transcription factor Rht1 - <i>Saccharomyces cerevisiae</i>	2e-85	SE7292	probable membrane protein YOR304a - yeast (<i>Saccharomyces cerevisiae</i>)	343.2	374.4	52.9	98.1	2e-25	01.01.04 phosphate metabolism
Pc13g19170	similarity to RNA polymerase II transcription factor Ugo2 - <i>Saccharomyces cerevisiae</i>	6e-06	NC0233440.1	gene: "B0D12.200", product: "conserved hypothetical protein", <i>Aspergillus niger</i>	189.6	189.6	12.0	12.0	1e-67	03.03.01 mitotic cell cycle and cell cycle control
Pc23g02260	similarity to transcription activator of lysine pathway Lys14 - <i>Saccharomyces cerevisiae</i>	0.0	T47118	hypothetical protein ZH2-Cy938 zinc-finger protein - fission yeast (<i>Schizosaccharomyces pombe</i>)	89.4	80.3	42.5	33.7	7e-36	01.03.01 purine nucleotide metabolism
Pc13g20830	strong similarity to protein Sot1 - <i>Saccharomyces cerevisiae</i>	1e-155	T49318	probable SOT1 protein [imported] - <i>Neurospora crassa</i>	191.1	183.9	134.4	72.6	0.0	03.03.01 mitotic cell cycle and cell cycle

Pc21g07940	strong similarity to transcription factor like protein An14g04590 - Aspergillus niger	16-79	AN15365.1	unannoted ORF. Sequence 22 from Patent WO02059324.	171.8	142.0	85.8	66.3	0	
Pc21g1091	strong similarity to zinc-finger transcription factor like protein An10g0800 - Aspergillus niger	26-21	NCB19418_21	gene: "B19H16.2307; product: "hypothetical protein";	231.0	259.0	220.0	118.0	26-34	05. PROTEIN SYNTHESIS
Pc21g1320	weak similarity to transcription activator of lysine pathway Lys14 - Saccharomyces cerevisiae	16-15	NC655_17	gene: "S55.1507; product: "hypothetical protein";	40.0	39.0	22.0	36.0	26-35	01.01.04. regulation of amino acid metabolism
Pc21g02940	similarity to hypothetical spliceosome associated protein Spf01 - Homo sapiens	36-52	AF42142.5	gene: "A1A1242.056; product: "hypothetical DNA] domain o	142.9	133.4	81.0	65.0	0	04.05. mRNA transcription
Pc21g1680	strong similarity to hypothetical C0 hypothetical protein CACD37.6.1 - Aspergillus fumigatus	16-47	AF42142.6	gene: "A1A1242.6; product: "hypothetical protein";	71.9	69.0	42.0	16.0	26-28	29. TRANSPOSABLE ELEMENTS, VIRAL AND PLASMID PROTEINS
Pc20g2150	strong similarity to DNA RNA helicase Hm1 - Saccharomyces cerevisiae	16-10	S61610	gene: "Hm1; product: "yeast [Saccharomyces cerevisiae]	31.4	34.1	24.5	16.0	17-15	0
Pc22g03450	weak similarity to U1 snRNP protein - Homo sapiens	26-19	NC12F1_22	gene: "12F1.21; product: "related to U1 SMALL NUCLEAR	95.1	97.2	40.0	30.0	94-46	03. CELL CYCLE AND DNA PROCESSING
Pc22g13510	weak similarity to 40 kD subunit of DNA-directed RNA polymerase I Rpa49 - Saccharomyces cerevisiae	46-08	T36336	probable DNA-directed RNA polymerase (EC 2.7.7.6) 149k d	68.2	69.2	63.0	30.1	1e-41	05.07. translational control
Pc22g13520	strong similarity to hypothetical transcription factor RAB4728.1 - Aspergillus oryzae	16-131	AB020371.1	product: "putative transcriptional activator"; Aspergillus nidula	31.4	34.1	24.5	16.0	17-15	04. TRANSCRIPTION
Pc22g14170	strong similarity to RNA Mat processing protein Gcd14 - Saccharomyces cerevisiae	26-96	NC2109_6	gene: "Y109.070; product: "related to GCN4 translational re	46.0	38.4	35.2	16.4	1e-162	03.03.01. mitotic cell cycle and cell cycle control
Pc22g16080	strong similarity to essential nucleolar protein Nop77 - Saccharomyces cerevisiae	16-101	T40151	ribosomal processing, ma binding, nucleolar pro tein - fission	150.1	130.1	128.0	51.0	0	04.01.01. RNA synthesis
Pc22g19520	strong similarity to transcription activator Adc2 - Saccharomyces cerevisiae	16-107	BX060808_2	gene: "G21B4.047; product: "related to transcription adaptor	254.9	260.4	104.7	14.77	0	40.10. nucleic
Pc22g21540	strong similarity to RNA polymerase I transcription factor RRN3 - Homo sapiens	36-86	BX42627_22	gene: "BB4227.22; product: "related to RNA Polymerase I Tr	256.9	215.8	157.3	91.0	0	10.01.01. unspecified signal transduction
Pc22g19510	strong similarity to XRN like protein An04g05060 - Aspergillus niger	36-31	NCB11823_15	gene: "B11823.1507; product: "conserved hypothetical prote	30.3	30.0	12.0	18.0	0	10.01.01. unspecified signal transduction
Pc24g11560	weak similarity to retrotransposon Tof1 - Nicotiana tabacum	16-132	T02206	hypothetical protein - common tobacco retrotransposon	44.7	40.7	33.1	16-109	0	01.04. phosphate metabolism
Pc24g19130	weak similarity to retrotransposon Tof1 - Nicotiana tabacum	16-136	T02206	hypothetical protein - common tobacco retrotransposon	298.7	940.7	164.8	136.71	1e-171	04.03.01. RNA synthesis
Pc13g16540	strong similarity to cytosolic aspartate--RNA ligase Dps1 - Saccharomyces cerevisiae	16-147	SVBYDC	aspartate-RNA ligase (EC 6.1.1.12), cytosolic [validated]; -y	137.11	1594.0	734.8	763.0	0	05.04. translation
Pc13g0380	strong similarity to glutamine--RNA ligase - Saccharomyces cerevisiae	0.0	SVBYCT	glutamine-RNA ligase (EC 6.1.1.19) - yeast [Saccharomyces	159.6	176.5	130.8	73.8	1e-157	04.05.01.04. translational control
Pc13g07040	strong similarity to translation initiation factor eIF-3 subunit like protein An14g01030 - Aspergillus niger	16-159	T38786	translation initiation factor eIF-3 - fission yeast [Schizosaccha	522.1	550.0	380.9	239.5	1e-100	05.04. translation
Pc16g02320	strong similarity to conserved eukaryotic protein involved in ribosome biogenesis Ebf1 - Saccharomyces cerevisiae	0.0	SS5459	probable membrane protein YMR0436 - yeast [Saccharomy	231.1	180.0	154.8	76.0	1e-116	01.03.16.01. RNA degradation
Pc16g0559	strong similarity to eukaryotic initiation factor E1F1 subunit TP11 - Ascidiposa italiana	46-58	OSJ003130_13	gene: "OSJ003009V.15; product: "Oryza sativa genomic DNA, c	317.6	381.1	207.6	125.4	0	05.04. translation
Pc16g09300	strong similarity to poly(ADP-ribose)-binding protein Pabp - Saccharomyces cerevisiae	0.0	ENU70731.1	gene: "b8M"; product: "putative poly(ADP-ribose) protein FaaM	490.4	438.0	422.4	186.1	1e-103	03.03.01. mitotic cell cycle and cell cycle control
Pc16g10350	strong similarity to translation initiation factor eIF-3 - Schizosaccharomyces pombe	0.0	T38716	translation initiation factor eIF-3 subunit [imputed] - fission ye	45.7	49.9	35.7	19.0	0	05.04. translation
Pc20g08010	strong similarity to translation initiation factor 3 complex protein like protein An11g10630 - Aspergillus niger	16-141	T40476	hypothetical protein SPBC464.04 - fission yeast [Schizosac	277.4	288.0	314.2	148.1	0	01.04. phosphate metabolism
Pc20g09480	strong similarity to valine--RNA ligase cpl-20 - Neurospora crassa	0.0	A41251	valine-RNA ligase (EC 6.1.1.9) - Neurospora crassa	359.7	407.0	271.8	186.2	3e-42	03.03.01. mitotic cell cycle and cell cycle control
Pc21g05120	strong similarity to poly(A)-specific ribonuclease Pan1 - Saccharomyces cerevisiae	96-65	S48440	poly(A)-specific ribonuclease (EC 3.1.13.4) - yeast [Sachar	99.8	85.3	68.8	37.0	7e-60	05.04. translation
Pc21g19420	strong similarity to mitochondrial ribosomal protein of the large subunit Yml35 - Saccharomyces cerevisiae	56-82	BX46220_10	gene: "B11E5.1007; product: "related to ribosomal protein Yml	175.2	189.9	18.8	789.9	2e-33	04.05.01.04. translational control
Pc21g21710	strong similarity to gamma subunit of translation elongation factor eEF-1 - Homo sapiens	36-36	AB056105.1	gene: "e1f1 gamma"; product: "elongation factor1 gamma"; C	1123.8	1703.9	941.1	670.1	1e-101	01.04. phosphate metabolism
Pc21g21020	strong similarity to nuclear protein gao2p - Schizosaccharomyces pombe	26-32	AF456933.1	product: "RNA binding protein"; Elaeis oleifera RNA binding	192.0	184.2	128.4	50.3	9e-94	06.01. protein folding and stabilization
Pc21g20100	strong similarity to protein kinase like protein An15g01310 - Aspergillus niger	0.0	AY456073.1	gene: "RHB15"; product: "putative response regulator receptor	360.0	328.0	167.6	114.9	5e-99	06.13. proteolytic degradation
Pc12g1120	strong similarity to protein precursor Bcl1 - Saccharomyces cerevisiae	66-114	T68717	protein BCL1 - fission yeast [imputed] - Neurospora cr	51.4	44.8	40.0	16.0	0	01.03.16.01. RNA degradation
Pc21g04620	strong similarity to aspartic protease Ypa3 - Saccharomyces cerevisiae	16-165	AB040478.1	gene: "opa3"; product: "aspartic protease"; Aspergillus oryza	51.8	50.0	20.3	11.3	0	06.07. protein modification
Pc21g03030	similarity to autophagy modulator Ap17 - Saccharomyces cerevisiae	46-14	S46993	probable membrane protein YBR217e - yeast [Saccharomy	271.4	248.0	120.0	15.0	0	06.07. protein modification
Pc12g1070	strong similarity to chaf-type molecular chaperone Yca1 - Aspergillus niger	76-10	T43723	chaf-type molecular chaperone beta [imputed] - Aspergillus	143.2	139.9	65.5	32.0	0	04.05.01.04. translation control
Pc12g12880	strong similarity to ubiquitin specific protease like protein An02g01420 - Aspergillus niger	16-87	S46936	hypothetical protein YAL197e - yeast [Saccharomyces cerevi	50.3	52.3	27.6	16.8	5e-48	04. TRANSCRIPTION
Pc21g13910	strong similarity to spermidine spermine N1-acetyltransferase like protein An14g03030 - Aspergillus niger	26-123	SPAC1002.7	gene: "asn1"; SPAC1002.07C; product: "N-acetyltransferase	145.6	136.8	48.8	36.0	5e-52	05. PROTEIN SYNTHESIS
Pc21g14240	strong similarity to GTPase like protein An14g02260 - Aspergillus niger	16-52	RYL2_1.10.0	gene: "RYL2_1.10.0; product: "putative GTPase-activating pr	219.0	214.0	89.0	87.2	0	06.01.01.04. translation control
Pc21g14810	strong similarity to zurin Ztn1 - Saccharomyces cerevisiae	16-123	S75072	probable zurin [imputed] - Neurospora crassa	426.8	413.5	368.4	206.4	7e-42	06.04. protein targeting, sorting and translocation
Pc21g16530	strong similarity to costomer gamma subunit 2 cosp2 - Homo sapiens	0.0	BX42668.1	gene: "B13D15.0507; product: "related to costomer gamma-2	1527.1	1289.2	415.1	313.5	5e-25	05. PROTEIN SYNTHESIS
Pc21g14510	strong similarity to SCF ubiquitin ligase Fbx1 - Saccharomyces cerevisiae	16-40	AK044953.1	gene: "Fbx1; product: "ubiquitin-protein ligase complexed	132.7	145.0	137.0	59.0	0	06.04. protein targeting, sorting and translocation
Pc21g15850	strong similarity to heat shock protein hsp30 - Saccharomyces nidulans	26-52	AB12668.1	gene: "hsp30"; product: "heat shock protein 30"; Aspergillu	35.1	29.4	22.9	12.3	2e-74	06.04. protein targeting, sorting and translocation
Pc24g0640	strong similarity to nuclear protein Eri1 - Saccharomyces cerevisiae	16-85	T39410	hypothetical protein SPBC19G1.09 - fission yeast [Schizosac	202.5	183.0	182.3	87.5	1e-150	06.04. protein targeting, sorting and translocation
Pc21g0400	strong similarity to nuclear protein Ndc80 - Saccharomyces cerevisiae	26-38	T40204	nuclear protein NDC80 - fission yeast [Schizosaccharomyces	200.3	190.0	124.0	59.0	0	06.04. protein targeting, sorting and translocation
Pc16g08000	strong similarity to mitochondrial import protein Mpi1 - Saccharomyces cerevisiae	16-149	AY121507.1	gene: "tim44"; product: "mitochondrial inner membrane trans	230.4	218.7	142.0	106.0	0	06.04. protein targeting, sorting and translocation
Pc16g11990	similarity to cdc25C associated protein kinase like protein An18g01800 - Aspergillus niger	46-09	G86927	protein HSB2E3.1 [imputed] - Caenorhabditis elegans	23.6	12.0	15.1	12.1	1e-174	06.04. protein targeting, sorting and translocation
Pc16g14760	strong similarity to karyopherin beta-1 - Saccharomyces cerevisiae	16-0	AY30543.1	gene: "kap61"; product: "putative karyopherin/importin beta-1"	269.0	269.0	173.0	178.0	0	0
Pc16g15440	weak similarity to ankrytin Ank2 - Homo sapiens	86-54	AS5575	ankyrin 3, long splice form - human	176.3	226.0	190.6	55.2	0	01.04. phosphate metabolism
Pc20g07070	similarity to receptor tyrosine kinase CKC-2 like protein An13g01500 - Aspergillus niger [putative pseudogene]	0.0			48.7	37.0	20.6	16.4	0	06.01. protein folding and stabilization
Pc20g19790	strong similarity to dual-specificity phosphatase mfp1 - Saccharomyces cerevisiae	16-26	T39517	0	209.8	186.0	106.0	66.0	1e-79	0
Pc20g09730	strong similarity to glutamyl-proline kinase like protein An12g0120 - Aspergillus niger	0.0		0	56.2	47.0	26.6	26.4	0	01.04.01.04. translational control
Pc20g10240	strong similarity to serine-type carboxypeptidase precursor cps5 - Aspergillus phoenicis	0.0	S78072	serine-type carboxypeptidase (EC 3.4.16.1) - Aspergillus nig	139.1	146.0	57.5	17.0	0	03. CELL CYCLE AND DNA PROCESSING
Pc20g11320	weak similarity to ankrytin 2 Ank2 - Drosophila melanogaster	36-11	AE017257_215	protein ankrytin repeat domain protein"; Wolbachia endosymb	12.8	12.8	10.0	4.0	0	06.04. protein targeting, sorting and translocation
Pc20g11430	strong similarity to protein required for accurate mitotic chromosome segregation Cnt1 - Saccharomyces cerevisiae	0.0	NCB18417.1	gene: "CNT1; product: "probable mitotic chromosome export	460.2	436.1	303.6	201.1	3e-42	04. TRANSCRIPTION
Pc20g10310	strong similarity to RAN-binding protein Kap123 - Saccharomyces cerevisiae	16-167	T39449	probable import beta-1 subunit - fission yeast [Schizosaccha	792.5	591.1	537.7	264.2	1e-120	01.04. phosphate metabolism
Pc21g09710	strong similarity to ADP-ribosylation factor-like protein Arf1 - Saccharomyces cerevisiae	66-29	AK009226.1	gene: "ARF1; product: "small GTP-binding protein"; Homo sap	48.8	25.5	18.2	12.8	2e-6	01.06.13. lipid and fatty-acid transport
Pc21g14860	strong similarity to seven kinase - Dictyostelium discoideum	16-157	NCB18417.1	gene: "B18M17.0507; product: "related to B18C20-like kinase	82.2	68.0	60.0	50.3	0	04.05.01.04. translation control
Pc21g16820	strong similarity to phosphatidylinositol-phosphatidylcholine transfer protein SEC14 - Yarrowia lipolytica	46-29	BX060899.19	gene: "B18C24.1907; product: "probable phosphatidylinositol	260.6	315.7	168.7	150.8	4e-45	06.04. protein targeting, sorting and translocation
Pc21g24540	similarity to signal recognition particle Sec63 - Saccharomyces cerevisiae	16-27	JG5816	signal recognition particle protein Sec63 - yeast [Yarrowia lip	261.6	254.3	127.7	134.0	2e-38	06.04. protein targeting, sorting and translocation
Pc20g0070	strong similarity to protein coat assembly protein MPS3 - Homo sapiens	66-57	AK042596.1	gene: "MPS3; product: "RNA binding protein"; Homo sapien	130.2	124.0	61.3	61.4	0	05. PROTEIN SYNTHESIS
Pc22g07060	similarity to vSNAARE Vcl1 - Saccharomyces cerevisiae	16-24	T40349	vesicle transport vSNAARE protein - fission yeast [Schizosaccha	425.4	415.4	251.7	194.3	1e-112	01.06.13. vesicular transport
Pc22g13560	strong similarity to vacuolar aminopeptidase Yac1 - Saccharomyces cerevisiae	16-170	NCB15020.13	gene: "B13D20.1607; product: "related to aminopeptidase yac	815.1	798.1	404.3	392.2	1e-157	01.06.13. fatty-acid and isoprenoid metabolism
Pc12g14430	strong similarity to vacuolar H(+)-Ca(2+) exchanger Yxc1 - Saccharomyces cerevisiae	16-40	AF103229.1	gene: "YXC1; product: "vacuolar H(+)-Ca(2+) exchanger"; Schiz	259.5	259.5	144.4	185.7	0	01.06.13. fatty-acid and isoprenoid metabolism
Pc21g16330	strong similarity to ABC-type vacuolar membrane protein Hmt1p - Schizosaccharomyces pombe	16-139	S25198	vacuolar membrane protein HMT1 - fission yeast [Schizosac	91.7	101.7	53.6	37.7	3e-75	01.01. amino acid metabolism
Pc15g13010	strong similarity to tetraacylin resistance protein tetA - Escherichia coli	16-11	AB037101	tetracycline resistance protein [imputed] - Agrobacterium tun	37.0	12.0	12.0	12.0	2e-54	01.05.07. C-compound, carbohydrate transport
Pc15g13630	strong similarity to choline permease Hmt1 - Saccharomyces cerevisiae	46-41	BX464968_108	gene: "hmtA"; product: "choline permease"; putative"; Aspergill	54.2	54.2	22.0	16.0	0	01.04.01.04. TRANSPORT AND TRANSPORT MECHANISMS
Pc15g12540	strong similarity to hypothetical transport protein YOR202c - Saccharomyces cerevisiae	0.0	BX469007.29	product: "possible transporter-like protein"; Aspergillus fumig	51.6	36.0	19.0	17.5	3e-43	08.19. cellular import
Pc15g16410	strong similarity to ABC-type vacuolar membrane protein hmt1p - Schizosaccharomyces pombe	0.0	S25198	vacuolar membrane protein HMT1 - fission yeast [Schizosac	1002.1	1034.4	716.0	400.7	1e-52	01.05.07. C-compound, carbohydrate transport
Pc15g19180	strong similarity to high affinity calcium transporter - Schizosaccharomyces pombe	16-39	PAN30117.1	gene: "YLP9"; product: "high affinity calcium transporter"; Sch	109.2	105.8	48.5	27.6	1e-161	05.09. other intracellular transport activities
Pc15g10660	strong similarity to membrane protein Pab1 - Saccharomyces cerevisiae	46-90	BX469007.11	gene: "mbr1"; product: "mbr-family multidrug resistance pro	55.9	38.1	22.6	13.9	4e-79	03.03.01.01. mitotic cell cycle
Pc21g07720	strong similarity to calcium-transporting ATPase Pmc1 - Saccharomyces cerevisiae	0.0	NCB19417.31	gene: "B19A17.3107; product: "putative calcium p-type ATPa	293.9	229.9	256.7	143.9	2e-57	04. TRANSCRIPTION
Pc21g17990	weak similarity to kinase K212b - Mus musculus	0.0	AY220423.1	gene: "K212b"; product: "kinase"; Bos taurus [validated]; -m	319.4	319.4	186.0	86.0	0	01.01.01.04. translation control
Pc21g11550	strong similarity to GTP-binding protein Rab4 - Homo sapiens	36-47	T38300	gene: "rab4"; product: "GTP-binding cassette multidrug trans	177.1	125.1	75.2	76.5	0	01.06.13. fatty-acid and isoprenoid metabolism
Pc21g13580	strong similarity to peptide transport protein Pab1 - Saccharomyces cerevisiae	0.0	AY187281.1	product: "d1m peptide transporter Z7; Phaeosphaeria nodorum	151.4	289.9	84.4	99.0	0	01.06.13. fatty-acid and isoprenoid metabolism
Pc21g13720	ABC transporter sequence asst from patient WO02013290-A2 - Penicillium chrysogenum	0.0	AF071410.1	gene: "atcC"; product: "ATP-binding cassette multidrug trans	84.6	80.0	29.1	4e-96	0	13.01.01. ion homeostasis
Pc21g13720	Pc21g13720 ABC transporter sequence asst from patient WO02013290-A2 - Penicillium chrysogenum	0.0	AF071410.1	gene: "atcC"; product: "ATP-binding cassette multidrug trans	84.6	80.0	29.1	4e-96	0	01.06.13. fatty-acid and isoprenoid metabolism
Pc21g19880	strong similarity to Na/H (+) antiporter ZSD02-22 - Zygosaccharomyces roullii	0.0	AY010962.2	gene: "ZSD9-2; product: "NhaP1"; Aspergillus fumigatus Mlp	147.3	166.1	111.1	67.1	4e-49	01.05.07. C-compound, carbohydrate transport
Pc21g01610	strong similarity to ABC-type vacuolar membrane protein hmt1p - Schizosaccharomyces pombe	0.0	S25198	vacuolar membrane protein HMT1 - fission yeast [Schizosac	759.9	823.1	492.6	403.0	1e-134	67.04. ion transporters
Pc21g0660	strong similarity to hypothetical amine transporter SPBC14.02 - Schizosaccharomyces pombe	16-53	T41145	product: "possible amine transporter - fission yeast [Schizosac	41.0	44.0	27.3	24.0	0	04.04.03.03. phenome response, mating-type determination, sex-specific proteins
Pc21g2380	strong similarity to mitochondrial tricarboxylate carrier - Rattus sp.	36-94	S67168	probable membrane protein YOR272c - yeast [Saccharomy	318.7	292.3	202.0	131.7	1e-105	08.13. vesicular transport
Pc21g27470	strong similarity to oligopeptide transporter Opl1 - Saccharomyces cerevisiae	16-133	T39467	hypothetical protein SPAC28B12.10c - fission yeast [Schiz	56.2	60.8	37.4	31.7	6e-47	40.03. cytoplasm

Pc13g0030	weak similarity to hypothetical protein An01g07930 - Aspergillus niger	0	0	0	158.3	85.1	69.5	61.8	0	
Pc13g0280	hypothetical protein	0	0	0	118.3	82.3	55.6	39.5	0	
Pc13g0840	strong similarity to hypothetical protein nc00832.1 - Neurospora crassa	0	0	0	64.5	54.1	58.1	27.5	0	
Pc13g0730	strong similarity to hypothetical protein contig5_part_1.1ta_1400wg - Aspergillus fumigatus	0	0	0	116.0	167.7	77.4	30.3	6e-80	14.04 cell differentiation
Pc13g0620	strong similarity to hypothetical protein contig1_part_1.153_scaffold_12.1ta_240wg - Aspergillus nidulans	0	0	0	98.8	80.7	85.2	30.2	0	99 UNCLASSIFIED PROTEINS
Pc13g0980	weak similarity to hypothetical protein BAB5353.1 - Homo sapiens	1e-08	BX897677_20	gene: "B15B10.180"; product: "hypothetical protein";	333.2	481.5	325.6	224.2	2e-38	11.05.05 virulence, disease factors
Pc13g01470	similarity to hypothetical protein SPAC12G12.02 - Schizosaccharomyces pombe	7e-11	T37599	hypothetical protein SPAC12G12.02 - fission yeast (Schizosaccharomyces pombe)	134.2	128.4	106.7	50.1	0	
Pc13g01640	weak similarity to hypothetical protein SPAC2AC3.05c - Schizosaccharomyces pombe	7e-63	T38346	hypothetical protein SPAC2AC3.05c - fission yeast (Schizosaccharomyces pombe)	326.8	354.4	322.5	246.2	2e-43	01.03 nucleotide metabolism
Pc13g02070	strong similarity to hypothetical protein contig1_2.96_scaffold_2.1ta_460wg - Aspergillus nidulans	0	0	0	470.9	171.0	136.3	124.4	1e-30	01.04 phosphate metabolism
Pc13g01910	strong similarity to hypothetical UDP-glucose 4-epimerase - Mesorhizobium loti	2e-78	B97596	UDP-glucose 4-epimerase [imported] - Aspergillus fumigatus	74.8	44.4	35.2	42.4	0	
Pc13g02380	strong similarity to hypothetical protein contig1_1.68_scaffold_1.1ta_200wg - Aspergillus nidulans	3e-18	BX842624_9	gene: "B10D6.090"; product: "related to putative cytoplasmic protein"	66.6	84.4	40.3	23.1	0	
Pc13g03200	strong similarity to hypothetical protein contig1_2.304wg - Aspergillus niger	0	0	0	99.9	80.2	49.2	615.0	0	10.01.99 other intracellular signal transduction activities
Pc13g03040	hypothetical protein	0	0	0	82.6	85.1	33.9	34.3	0	
Pc13g03430	strong similarity to hypothetical protein Kex3 - Saccharomyces cerevisiae	0.0	S55151	probable membrane protein YNL120w - yeast (Saccharomyces cerevisiae)	173.0	153.4	97.7	58.9	1e-40	08.13 vacuolar transport
Pc13g03450	strong similarity to hypothetical mixed-linked glucanase precursor related protein MLG1 - Neurospora crassa	1e-72	U91606_1	gene: "MLG1"; product: "mixed-linked glucanase precursor";	63.8	10.1	12.0	12.0	0	
Pc13g03810	strong similarity to hypothetical protein contig31_part_1.1ta_890wg - Aspergillus fumigatus	2e-20	T40913	probable integral membrane protein - fission yeast (Schizosaccharomyces pombe)	124.9	84.4	59.1	48.2	0	
Pc13g03880	strong similarity to hypothetical protein An12g04010 - Aspergillus niger	3e-04	AF133896_1	gene: "biga"; product: "putative surface-exposed virulence protein"	419.6	394.0	282.6	191.0	1e-08	
Pc13g04190	strong similarity to hypothetical protein of the pastille biosynthesis gene cluster paxU - Penicillium parvii	1e-123	AF27868_7	gene: "paxU"; product: "Penicillium parvii dimethylallyl pyrophosphate synthase"	29.1	12.1	12.3	13.9	1e-99	06.04 protein targeting, sorting and translocation
Pc13g04460	strong similarity to hypothetical protein contig1_1.88_scaffold_1.1ta_690wg - Aspergillus nidulans	2e-07	BX842638_16	gene: "G1767.160"; product: "putative protein";	153.2	138.1	83.2	54.4	0	
Pc13g04790	strong similarity to hypothetical protein CG3196-PB - Drosophila melanogaster	2e-48	AE035376_35	product: "CG3196-PB"; Drosophila melanogaster choromosome 3	49.2	44.4	36.1	16.9	0	
Pc13g04830	similarity to hypothetical protein An02g06800 - Aspergillus niger	0	0	0	23.8	24.1	12.1	12.0	0	
Pc13g07350	strong similarity to hypothetical protein contig1492_0.1ta_6190wg - Aspergillus fumigatus	1e-06	AY281368_1	gene: "cip2"; product: "Cip2"; Hypocrea jecorina strain OM66	18.5	47.5	43.6	14.3	0	01.05 C-compound and carbohydrate metabolism
Pc13g07600	strong similarity to hypothetical protein contig1488_2.1ta_850wg - Aspergillus fumigatus	2e-04	AY164486_1	gene: "Pkd113"; product: "polycystic kidney disease 1-like 3"	58.7	51.6	31.7	18.9	0	
Pc13g08140	strong similarity to hypothetical protein contig1486_2.1ta_1050wg - Aspergillus fumigatus	4e-13	NC89K_17_10	gene: "B9K17.090"; product: "conserved hypothetical protein"	128.0	45.1	88.6	59.1	0	
Pc13g08150	similarity to hypothetical protein An04g03190 - Aspergillus niger	0	0	0	349.8	128.0	219.4	139.4	0	
Pc13g08160	hypothetical protein	0	0	0	80.6	42.1	56.8	26.5	0	
Pc13g08010	strong similarity to hypothetical protein An02g07410 - Aspergillus niger	0	0	0	280.7	349.0	146.5	98.9	0	
Pc13g09000	strong similarity to hypothetical protein contig336.1ta_610wg - Aspergillus fumigatus	1e-04	AB083477_1	gene: "Pico1"; product: "Pico1"; Mus musculus Pico mRNA	103.1	42.0	50.1	50.7	0	
Pc13g09280	strong similarity to hypothetical protein An02g06790 - Aspergillus niger	2e-12	CN509545_42	DNA centromeric region sequence from BAC DP26B06, DP3	18.1	37.3	12.0	12.9	1e-94	14.04 cell differentiation
Pc13g09400	strong similarity to hypothetical protein An02g09800 - Aspergillus niger	2e-14	CN509545_56	DNA centromeric region sequence from BAC DP26B06, DP3	40.4	40.4	19.2	19.9	0	
Pc13g09820	similarity to hypothetical protein F17H10.1 - Caenorhabditis elegans	1e-17	T21079	hypothetical protein F17H10.1 - Caenorhabditis elegans	786.3	456.5	214.1	142.0	0	
Pc13g10170	weak similarity to hypothetical protein BB24.160 - Neurospora crassa	8e-04	AF345640_1	gene: "Myb3a1"; product: "myb-binding protein 1a";	60.7	35.3	31.9	16.7	0	
Pc13g10220	strong similarity to hypothetical protein An15g0870 - Aspergillus niger	0	0	0	132.5	150.6	72.7	44.1	0	
Pc13g10380	strong similarity to hypothetical protein An16g0440 - Aspergillus niger	0	0	0	67.2	44.4	20.1	16.7	0	
Pc13g10610	strong similarity to hypothetical protein An16g0380 - Aspergillus niger	8e-45	BX08811_2	gene: "H4H7.020"; product: "putative protein";	245.4	236.5	261.0	116.8	0	
Pc13g10720	hypothetical protein	0	0	0	171.8	173.1	115.2	68.7	0	
Pc13g10840	strong similarity to hypothetical transcription factor Ite-like protein CAE4709.1 - Aspergillus fumigatus	1e-117	BX849005_80	product: "possible transcription factor Ite-like protein"; Aspergillus fumigatus	176.5	82.5	40.4	40.1	0	
Pc13g10980	strong similarity to hypothetical protein An08g03680 - Aspergillus niger	4e-45	BX849005_48	product: "hypothetical protein, putative"; Aspergillus fumigatus	287.0	217.4	101.8	88.6	0	
Pc13g11260	strong similarity to hypothetical protein contig1_1.51_scaffold_1.1ta_100wg - Aspergillus nidulans	2e-05	AP029596_159	gene: "mlt105"; Mesorhizobium loti DNA, complete genome	26.8	12.0	12.0	12.0	0	
Pc13g11380	strong similarity to hypothetical protein contig1_1.1ta_100wg - Aspergillus fumigatus	6e-25	BX849008_23	product: "hypothetical protein"; Aspergillus fumigatus BAC pil	49.1	55.2	44.1	25.5	0	
Pc13g11610	strong similarity to hypothetical protein An01g1780 - Aspergillus niger	5e-06	BX71859_231	Photobacterium luminescens subsp. laumondii T101 complete genome	53.1	54.2	28.1	20.2	1e-40	98 CLASSIFICATION NOT YET CLEAR-CUT
Pc13g11620	strong similarity to hypothetical protein - Aspergillus fumigatus	1e-114	BX849006_30	product: "hypothetical protein"; Aspergillus fumigatus BAC pil	93.5	41.6	64.5	63.1	0	
Pc13g12370	strong similarity to hypothetical protein YLR003b - Saccharomyces cerevisiae	1e-107	BX849006_37	product: "hypothetical protein"; Aspergillus fumigatus BAC pil	56.9	65.5	44.1	49.9	0	
Pc13g13090	hypothetical protein	0	0	0	118.1	133.8	84.3	57.6	0	
Pc13g13230	strong similarity to hypothetical protein B9H12.00 - Neurospora crassa	1e-76	BX849005_117	product: "hypothetical protein, conserved"; Aspergillus fumigatus	165.0	173.3	198.0	88.9	3e-31	01.04 phosphate metabolism
Pc13g13310	hypothetical protein	0	0	0	118.1	133.8	84.3	57.6	0	
Pc13g13430	similarity to serine-repeat antigen 4 - Plasmodium vivax	2e-14	T30841	serine-repeat antigen 4 - Plasmodium vivax	55.3	46.0	29.0	12.0	0	
Pc13g13670	strong similarity to hypothetical protein SPAC08B.07 - Schizosaccharomyces pombe	3e-21	T40768	hypothetical protein SPAC08B.07 - fission yeast (Schizosaccharomyces pombe)	79.3	89.8	84.7	41.6	3e-39	01.05.01.01 sugar, polyside, glycoly and carboxylate catabolism
Pc13g14080	similarity to hypothetical protein An02g01812.10 - Aspergillus niger	2e-07	NC201810_17_0	gene: "B10A7.100"; product: "hypothetical protein";	42.7	102.8	110.1	55.2	0	
Pc13g14090	strong similarity to hypothetical suppressor of bent1 bud5 mutant - Schizosaccharomyces pombe	1e-19	CG202138_1	gene: "63305GSM11R1K"; product: "63305GSM11R1K protein";	158.4	130.4	61.3	132.1	1e-60	20.25.01.03 hearing
Pc13g14120	strong similarity to hypothetical protein An02g09000 - Aspergillus niger	0	0	0	441.6	399.8	325.4	150.1	0	
Pc13g14140	similarity to hypothetical protein An02g08010 - Aspergillus niger	4e-20	AE035376_35	product: "CG3196-PB"; Drosophila melanogaster choromosome 3	49.2	44.4	36.1	16.9	0	
Pc13g14200	strong similarity to hypothetical protein contig61.1ta_380wg - Aspergillus fumigatus	6e-45	T45696	related to nlf-specific regulatory protein [imported] - Neurospora crassa	74.5	41.4	58.9	100.0	1e-113	99 UNCLASSIFIED PROTEINS
Pc13g14240	similarity to hypothetical protein An02g09070 - Aspergillus niger	0	0	0	216.2	294.3	168.8	92.2	0	
Pc13g14470	strong similarity to hypothetical protein YOR311c - Saccharomyces cerevisiae	1e-91	NC89B_11_21	gene: "B9B11.210"; product: "conserved hypothetical protein"	288.0	271.4	182.0	134.5	0	
Pc13g14490	weak similarity to hypothetical protein contig1.1ta_880wg - Aspergillus fumigatus	0	0	0	140.2	136.5	72.2	43.8	0	
Pc13g15240	strong similarity to hypothetical protein contig61.1ta_107wg - Aspergillus fumigatus	2e-12	NC05543_5	gene: "G6SA3.060"; product: "hypothetical protein";	191.8	82.5	35.1	21.2	0	
Pc13g15250	strong similarity to hypothetical transcriptional activator ptaA - Aspergillus nidulans	8e-37	AF205977_1	gene: "ptaA"; product: "PTAA"; Aspergillus nidulans PTAA (g)	39.1	86.2	23.3	16.0	0.0	03.01.03 DNA synthesis and replication
Pc13g15370	strong similarity to hypothetical protein contig51.1ta_1380wg - Aspergillus fumigatus	0	0	0	349.3	292.3	223.3	49.5	0	
Pc13g15480	strong similarity to hypothetical DNA replication licensing factor rimQ - Aspergillus nidulans	0.0	AF014813_1	gene: "rimQ"; product: "DNA replication licensing factor";	276.8	174.0	134.3	104.8	0	
Pc13g15640	weak similarity to hypothetical protein RK272 - Mycobacterium tuberculosis	4e-09	AE017240_191	product: "hypothetical protein"; Mycobacterium avium subsp.	162.7	47.1	86.0	48.5	2e-34	01.04 phosphate transport
Pc13g15660	strong similarity to hypothetical protein contig1.1ta_1610wg - Aspergillus fumigatus	0	0	0	159.8	174.8	122.4	74.8	0	99 UNCLASSIFIED PROTEINS
Pc14g01600	similarity to hypothetical monooxygenase Eabp6 - Saccharomyces cerevisiae	3e-20	AE030696_41	gene: "kar"; product: "CG1228B-PA"; Drosophila melanogaster	30.0	12.0	13.5	12.0	2e-27	04.05.01.04 transmembrane control
Pc14g01710	strong similarity to hypothetical protein contig1480_3.1ta_530wg - Aspergillus fumigatus	2e-82	AE052974_1	unnammed ORF; Sequence 2 from Patent WO030397025.	35.2	122.0	12.0	12.0	1e-73	14.04 cell differentiation
Pc14g01720	strong similarity to hypothetical protein contig1480_3.1ta_530wg - Aspergillus fumigatus	4e-09	AY327407_1	product: "hypothetical protein"; Homo sapiens hypothetical p	173.6	161.8	123.4	84.6	4e-53	99 UNCLASSIFIED PROTEINS
Pc14g05850	strong similarity to hypothetical protein CG135 - Homo sapiens	6e-61	CN5L1T196_1	unnammed ORF; human full-length cDNA clone C50D020Y11	273.1	332.4	203.0	146.3	1e-100	63.03 nucleic acid binding
Pc14g06890	strong similarity to hypothetical conserved protein SPAC13A2.02 - Schizosaccharomyces pombe	4e-38	NC18F11_16	gene: "18F11.160"; product: "conserved hypothetical protein";	119.8	118.1	96.2	47.6	0	
Pc14g12200	strong similarity to hypothetical protein YOR551c - Saccharomyces cerevisiae	1e-43	SC1328353_49	gene: "YOR551c"; S. cerevisiae YOR551c DNA fragment	439.4	385.5	193.4	169.7	3e-74	99 UNCLASSIFIED PROTEINS
Pc14g1480	similarity to hypothetical protein An16g06430 - Aspergillus niger	7e-04	AE014753_3	product: "hypothetical protein"; Bifidobacterium longum NCC2	529.6	385.0	291.9	144.2	3e-35	06.07 protein modification
Pc14g14940	weak similarity to hypothetical protein YNL051w - Saccharomyces cerevisiae	2e-58	NC82F7_3	gene: "B2F7.030"; product: "conserved hypothetical protein";	149.8	144.3	67.0	32.9	3e-49	04.05.01.04 transmembrane control
Pc14g15450	strong similarity to hypothetical conserved protein encoded by B2F7.170 - Neurospora crassa	1e-24	NC82F7_15	gene: "B2F7.170"; product: "conserved hypothetical protein";	205.9	161.1	74.3	33.7	4e-63	99 UNCLASSIFIED PROTEINS
Pc14g17130	strong similarity to hypothetical protein contig1_1.98_scaffold_1.1ta_1210wg - Aspergillus nidulans	7e-04	BC055247_1	gene: "zgc-63801"; product: "similar to eukaryotic translation product";	730.6	790.8	472.1	370.6	0	
Pc14g1850	strong similarity to hypothetical protein contig1482_0.1ta_440wg - Aspergillus fumigatus [putative sequencing error]	1e-36	AC116977_25	product: "similar to Mus musculus (Mouse). hypothetical prot	162.9	162.9	88.4	93.8	0	
Pc14g19570	strong similarity to hypothetical protein contig1482_0.1ta_520wg - Aspergillus fumigatus	1e-32	T51915	hypothetical protein B23H1.270 [imported] - Neurospora crassa	149.7	136.3	74.5	64.6	0	
Pc14g2100	hypothetical protein	0	0	0	69.6	62.3	77.0	32.7	0	
Pc14g2120	strong similarity to hypothetical protein An05g02110 - Aspergillus niger	0	0	0	21.8	24.4	12.0	12.0	0	
Pc15g0460	weak similarity to hypothetical protein contig127.1ta_110wg - Aspergillus fumigatus	0	0	0	43.6	47.8	38.2	163.1	0	
Pc15g0560	weak similarity to hypothetical protein SPCC045.13 - Schizosaccharomyces pombe	0	0	0	55.2	32.3	12.0	16.1	4e-53	01.03.16.01 RNA degradation
Pc15g1220	strong similarity to hypothetical protein contig1487_1.1ta_1750wg - Aspergillus fumigatus	0	0	0	384.5	277.9	165.5	89.7	0	
Pc15g1230	strong similarity to hypothetical protein ac2a01303091 - Kluyveromyces marxianus	8e-28	AF410318_1	product: "AT4g24770F22K16.30"; Arabidopsis thaliana AT4	413.6	407.8	382.2	163.1	1e-83	01.06.16. fatty-acid and isoprenoid metabolism
Pc15g1530	hypothetical protein	0	0	0	316.1	432.5	153.6	107.2	0	
Pc16g0080	strong similarity to lipase - Penicillium aili	1e-151	AY303124_1	gene: "lipA"; product: "lipase"; Penicillium aili lipase (lipA)	95.1	16.4	36.1	12.0	0	
Pc16g0970	similarity to hypothetical protein B14D6.5 - Neurospora crassa	3e-05	T45605	hypothetical protein B14D6.570 [imported] - Neurospora crassa	61.3	48.6	29.2	15.4	1e-78	99 UNCLASSIFIED PROTEINS
Pc16g1810	strong similarity to hypothetical protein contig16g_part_1.1ta_70wg - Aspergillus fumigatus	0	0	0	48.0	48.2	23.4	21.8	6e-65	
Pc16g1820	strong similarity to hypothetical protein An08g02080 - Aspergillus niger	1e-66	NC84N11_10	gene: "B24N11.100"; product: "conserved hypothetical protein"	153.3	175.8	119.4	86.7	1e-08	05.04 translation
Pc16g2370	similarity to hypothetical protein SPBC16B.02c - Schizosaccharomyces pombe	2e-15	T39677	hypothetical protein SPBC16B.02c - fission yeast (Schizosaccharomyces pombe)	367.3	334.4	187.4	137.0	2e-49	04 TRANSCRIPTION
Pc16g2480	strong similarity to hypothetical branched-chain alpha-ketoacid dehydrogenase E1 beta subunit - Homo sapiens	2e-97	T46720	translational initiation factor 1 (E1 beta subunit) related protein I	444.9	491.4	244.0	149.0	0	
Pc16g2560	strong similarity to hypothetical protein IAB.260 - Schizosaccharomyces pombe	3e-35	NC18A_15	gene: "IAB.220"; product: "conserved hypothetical protein";	355.3	341.4	232.9	105.5	0	
Pc16g2620	hypothetical protein	0	0	0	33.4	13.1	19.3	12.0	0	
Pc16g3130	strong similarity to hypothetical protein I190_scaffold_2.1ta_540wg - Fusarium graminearum	0	0	0	139.5	166.1	141.5	89.4	4e-92	1

Pc16g10730	similarity to hypothetical protein contig12.1ta_260wg - Aspergillus fumigatus	0	0	89.3	36.1	36.1	16.3	1e-47	03.03.01.01.11 mitosis
Pc16g11580	hypothetical protein	0	0	137.4	74.5	81.5	25.9	0	
Pc16g12330	similarity to hypothetical protein YNL24w - Saccharomyces cerevisiae	1e-101	BX64907.20	0	23.9	24.1	12.0	12.0	0
Pc16g12480	similarity to hypothetical protein AAM5689.1 - Xanthomonas axonopodis	1e-147	BX64907.31	0	36.9	36.9	42.7	24.9	0
Pc16g12590	strong similarity to hypothetical protein - Aspergillus fumigatus	4e-46	BX64907.38	0	76.7	76.7	88.3	14.1	4e-87
Pc16g13340	strong similarity to hypothetical protein contig46_part_1.1ta_780wg - Aspergillus fumigatus	1e-77	BX64907.41	0	30.7	30.5	12.0	12.0	0
Pc16g13080	strong similarity to hypothetical protein F12L6.11 - Arabidopsis thaliana	0.0	BX64907.82	0	194.2	201.2	111.6	91.8	0
Pc16g13380	similarity to hypothetical protein mg2520.1 - Magnaporthe oryzae	0	0	129.1	157.2	160.9	62.9	1e-15	01.04 phosphate metabolism
Pc16g13550	weak similarity to hypothetical protein - Streptomyces avermitilis	3e-09	AP050545.235	0	129.7	143.2	85.2	54.3	2e-46
Pc16g13600	strong similarity to hypothetical protein kinase Krd3 - Saccharomyces cerevisiae	0.0	BX42682.3	0	177.0	135.5	96.1	64.3	2e-81
Pc16g14160	strong similarity to hypothetical protein SM20006 - Sinorhizobium meliloti	2e-16	BX52696.171	0	197.9	207.1	162.9	101.9	0
Pc16g14620	strong similarity to hypothetical protein contig46_part_1.1ta_1500wg - Aspergillus fumigatus	2e-97	102038.15_17	0	144.1	141.4	69.3	101.4	0
Pc16g15040	strong similarity to hypothetical protein contig1.116_scaffold_8.1ta_60wg - Aspergillus nidulans	0	0	0	44.5	29.1	12.0	12.0	0
Pc16g15460	hypothetical protein	0	0	0	290.7	654.1	285.3	98.7	1e-136
Pc17g01090	weak similarity to hypothetical protein An02g01040 - Aspergillus niger [truncated ORF][putative pseudogene]	0	0	0	41.8	21.3	13.5	15.5	0
Pc17g00150	similarity to hypothetical transposase - Topolycodium infatum [truncated ORF][putative pseudogene]	3e-28	TRESTLES.1	gene: "trp"; product: "putative transposase"; T.infatum trans	156.7	141.7	112.3	62.7	0
Pc17g00210	hypothetical protein	0	0	0	31.8	12.0	12.0	12.0	0
Pc17g00420	hypothetical protein	0	0	0	91.0	94.5	37.5	31.5	0
Pc17g00680	hypothetical protein	0	0	0	24.6	30.5	19.6	12.0	0
Pc17g00780	hypothetical protein	1e-04	T31793	enamelin matrix protein - mouse	1868.7	1820.8	1248.3	651.9	0
Pc17g01180	hypothetical protein	0	0	0	31.8	23.3	12.0	12.0	0
Pc17g01240	weak similarity to hypothetical protein contig38.1ta_140wg - Aspergillus fumigatus [truncated ORF][putative sequencing error]	0	0	0	70.1	73.4	41.0	3e-75	01.07.01 biosynthesis of vitamins, cofactors, and prosthetic groups
Pc18g00880	strong similarity to hypothetical transcription regulator meabB - Aspergillus nidulans	1e-104	ANMEABG.1	gene: "meabB"; product: "MEAB protein"; Aspergillus nidulans	362.5	368.9	303.4	183.0	3e-36
Pc18g00980	strong similarity to hypothetical trunk lateral cell specific gene HTLC1 - Heterosyria rosea	0.0	NC23261.21	gene: "B2031.210"; product: "conserved hypothetical protein"	594.3	428.8	524.4	235.1	0
Pc18g01480	strong similarity to putative transcriptional regulator CAF32162.1 - Aspergillus fumigatus	3e-25	BX64907.85	gene: "putative transcriptional regulator, putative"; Aspergillus fum	33.9	35.8	14.1	16.0	4e-46
Pc18g01680	similarity to hypothetical protein An02g08040 - Aspergillus niger	0	0	0	36.6	29.4	12.0	12.0	0
Pc18g02200	strong similarity to hypothetical protein contig1.1_06_scaffold_4.1ta_140wg - Aspergillus nidulans	2e-06	AK01861.16	Mus musculus 16 days embryo head cDNA, RKEN full-length	137.5	61.8	42.3	16.7	0
Pc18g02760	strong similarity to hypothetical aldehyde dehydrogenase CAD36564.1 - Schizosaccharomyces pombe	1e-164	T50272	probable aldehyde dehydrogenase [imported] - fission yeast	482.1	313.1	141.7	179.1	2e-58
Pc18g03070	similarity to hypothetical protein SPAC6F8.17 - Schizosaccharomyces pombe	4e-36	AY04033.1	gene: "tap1"; product: "Tap1"; Schizosaccharomyces pombe	160.6	145.2	64.7	58.3	3e-38
Pc18g03270	strong similarity to spem tail-specific protein An1g05500 - Aspergillus niger	5e-05	S33760	hypothetical protein YKL062 - yeast [Saccharomyces cerev	157.3	168.3	86.3	43.9	0
Pc18g03440	strong similarity to hypothetical protein contig46_part_2.1ta_1500wg - Aspergillus fumigatus	6e-05	S14620	botosomal protein YN25, mitochondrial - yeast [Saccharomy	104.4	104.5	75.7	34.2	4e-62
Pc18g03770	weak similarity to hypothetical protein contig589.1ta_560wg - Aspergillus fumigatus	0	0	0	44.6	41.0	12.0	12.0	0
Pc18g03880	strong similarity to hypothetical protein An07g07410 - Aspergillus niger	7e-14	PFMAL13.585	gene: "MAL13P1.302"; product: "hypothetical protein"; Plas	113.4	106.8	64.5	41.7	1e-45
Pc18g03960	weak similarity to hypothetical protein HPRF03072 - Homo sapiens	0	0	0	127.2	127.6	54.1	79.9	0
Pc18g04150	strong similarity to hypothetical protein nc008637.1 - Neurospora crassa	3e-10	CEAF3131.1	gene: "unc-89"; product: "Uncoordinated protein 89, isoform B	89.0	105.5	63.2	33.3	0
Pc18g04160	strong similarity to polyprotein sequence like protein An07g07480 - Aspergillus niger [putative sequencing error]	1e-48	T38840	hypothetical tyrosine repeats containing protein - fission yeast	239.7	205.1	133.2	81.7	0
Pc18g04450	strong similarity to hypothetical protein contig1462.2.1ta_1500wg - Aspergillus fumigatus	3e-04	AK030464.1	unannotated ORF; Homo sapiens cDNA FLJ017615.1s, clone TE	183.4	183.4	143.8	74.9	0
Pc18g04570	strong similarity to hypothetical protein contig598.1ta_60wg - Aspergillus fumigatus	0	0	0	55.2	42.1	26.5	20.8	2e-44
Pc18g04810	strong similarity to hypothetical protein An07g07140 - Aspergillus niger	7e-04	AF350448.1	product: "enkephope glycoprotein precursor"; Cnimean-Congo h	195.0	149.3	58.5	50.0	2e-39
Pc18g04940	strong similarity to Yopep like protein An07g07170 - Aspergillus niger	2e-06	S57998	gene: "B2050664.1"; product: "hypothetical protein NC010507"; Homo sapiens	272.3	249.3	152.2	93.6	0
Pc18g05260	similarity to Ntp2 - Saccharomyces cerevisiae	4e-10	S57998	NEBP protein - yeast [Saccharomyces cerevisiae]	95.5	80.2	51.7	36.7	0
Pc18g05650	strong similarity to hypothetical protein An07g05260 - Aspergillus niger	0	0	0	104.8	80.5	22.8	24.1	0
Pc18g05810	strong similarity to hypothetical protein An07g05810 - Aspergillus niger	0	0	0	91.0	84.4	42.7	47.6	0
Pc18g05990	similarity to hypothetical protein SPAC26A3.14c - Schizosaccharomyces pombe	3e-04	T38402	hypothetical protein SPAC26A3.14c - fission yeast [Schizos	38.6	33.5	12.0	14.6	0
Pc18g06400	similarity to hypothetical protein An07g06260 - Aspergillus niger	7e-06	AF156773.22	gene: "ORF21"; product: "ORF21 protein"; Gibberella monili	59.4	65.0	43.1	29.6	0
Pc18g06610	hypothetical protein	0	0	0	169.7	164.7	80.7	16.7	0
Pc18g06590	hypothetical protein	0	0	0	134.8	106.7	29.7	28.2	3e-92
Pc18g06880	hypothetical protein	0	0	0	66.8	12.0	12.0	12.0	4e-73
Pc19g01070	similarity to hypothetical protein - Oryza sativa	3e-21	AP020371.11	Oryza sativa [japonica cultivar group] genomic DNA, chromo	29.8	48.1	12.0	12.0	0
Pc20g01160	strong similarity to hypothetical protein mg094979.1 - Magnaporthe oryzae	4e-44	SCMP3P.3	gene: "SMPP3"; product: "SMPP3 protein"; S.cerevisiae SMP3	46.3	30.1	25.2	14.8	0
Pc20g01260	similarity to hypothetical protein AAK58053.1 - Ophiostoma novo-umi	2e-14	AF378550.1	product: "unknown"; Ophiostoma novo-umi clone 13 unknow	79.3	92.1	32.8	26.4	0
Pc20g01380	strong similarity to hypothetical protein An07g01650 - Aspergillus niger	0	0	0	37.4	38.9	20.2	12.1	0
Pc20g01480	strong similarity to hypothetical protein An07g02040 - Aspergillus niger	0	0	0	69.3	67.1	31.1	17.0	0
Pc20g01510	strong similarity to hypothetical protein An06g05278 - Aspergillus niger	0	0	0	126.3	89.1	49.5	31.0	0
Pc20g01580	similarity to hypothetical protein An07g02120 - Aspergillus niger	0	0	0	118.7	87.5	47.5	2e-63	00.01 protein folding and isopren
Pc20g01590	strong similarity to hypothetical protein CAD21225.1 - Neurospora crassa	0.0	AY522343.1	product: "HrbB"; Emericella nidulans HrbB gene, compleme	186.7	134.7	90.5	81.0	0.0
Pc20g01600	similarity to hypothetical AAA-ATPase AAR34293.1 - Geobacter sulfurreducens	5e-17	AY02101.117	product: "ATPase, AAA family"; Geobacter sulfurreducens P	891.2	474.0	402.1	193.2	0
Pc20g01860	strong similarity to hypothetical protein contig42.1ta_690wg - Aspergillus fumigatus	1e-115	T52121	acyl-CoA oxidase (EC 1.3.3.6), peroxisomal [elaidated] - Anal	438.6	181.1	125.4	220.0	0
Pc20g01950	strong similarity to hypothetical protein contig1462.2.1ta_1500wg - Aspergillus fumigatus	6e-05	SCAR01GN.8	gene: "saHP7"; product: "saHP7 protein"; Saccharomyces mud	83.1	72.4	59.3	34.2	0
Pc20g02060	strong similarity to hypothetical protein An02g13370 - Aspergillus niger	9e-72	BX42638.30	gene: "TC14.100"; product: "hypothetical protein"; Neurosp	107.6	101.5	60.8	41.8	2e-59
Pc20g02150	strong similarity to hypothetical protein An02g13370 - Aspergillus niger	0	0	0	77.9	62.8	33.0	28.4	2e-55
Pc20g02780	strong similarity to hypothetical protein PA3829 - Pseudomonas aeruginosa	3e-60	BX3167	hypothetical protein PA3829 [imported] - Pseudomonas aeru	117.0	112.0	64.8	30.0	0
Pc20g02860	similarity to hypothetical myb-like DNA binding protein SPAC13G7.10 - Schizosaccharomyces pombe	0	0	0	31.1	17.5	12.0	12.0	0
Pc20g03030	strong similarity to hypothetical protein contig1487.1.1ta_1020wg - Aspergillus fumigatus	5e-12	BX42641.13	gene: "B2221.140"; product: "conserved hypothetical protein"	343.1	275.1	170.0	102.5	1e-105
Pc20g03190	strong similarity to hypothetical protein An11g07570 - Aspergillus niger	0	0	0	103.0	104.3	46.2	1e-34	40 SUBCELLULAR LOCALISATION
Pc20g03790	strong similarity to hypothetical protein SPBC1685.14c - Schizosaccharomyces pombe	0	0	0	297.2	277.0	207.5	133.3	0
Pc20g04000	similarity to hypothetical protein YOR252w - Saccharomyces cerevisiae	1e-12	S67149	hypothetical protein YOR252w - yeast [Saccharomyces cere	178.0	146.5	103.3	68.6	0
Pc20g04190	weak similarity to hypothetical protein SPAC6B12.11 - Schizosaccharomyces pombe	0	0	0	96.4	87.3	46.8	27.3	0
Pc20g04290	hypothetical protein	0	0	0	34.4	48.7	19.4	17.6	4e-53
Pc20g04630	strong similarity to hypothetical protein contig1.110_scaffold_8.1ta_780wg - Aspergillus nidulans	0	0	0	35.4	12.0	12.0	12.0	0
Pc20g04700	weak similarity to hypothetical protein F17H1.1 - Caenorhabditis elegans	7e-14	B97746	hypothetical protein RC0370 [imported]; Rickettsia corali	79.2	67.1	32.0	12.0	0
Pc20g05020	weak similarity to hypothetical coiled-coil protein - Schizosaccharomyces pombe	4e-29	NCB7423.23	gene: "B7H23.240"; product: "conserved hypothetical protein"	44.2	30.1	20.2	15.5	2e-30
Pc20g05210	similarity to hypothetical protein nc009167.1 - Neurospora crassa	0	0	0	48.5	22.8	25.3	12.0	2e-37
Pc20g05330	similarity to hypothetical mitochondrial ribosomal protein - Schizosaccharomyces pombe	1e-12	T41110	hypothetical protein SPCC1739.02c - fission yeast [Schizos	277.7	255.5	174.4	135.1	1e-39
Pc20g05480	weak similarity to hypothetical forkhead nuclear signaling protein SPBC3H7.13 - Schizosaccharomyces pombe	7e-12	NC065A3.6	gene: "G65A3.080"; product: "putative protein"; Neurospora c	819.7	662.3	337.1	256.7	0
Pc20g05610	strong similarity to hypothetical protein contig589.1ta_1510wg - Aspergillus fumigatus	3e-53	NC12F11.8	gene: "12F11.080"; product: "hypothetical protein"; Neurosp	68.9	75.4	33.5	32.7	0
Pc20g05970	similarity to hypothetical protein An02g12880 - Aspergillus niger	0	0	0	237.3	146.5	72.6	62.2	1e-41
Pc20g05970	similarity to hypothetical protein An02g12880 - Aspergillus niger	0	0	0	58.7	47.5	22.8	20.0	3e-79
Pc20g06000	strong similarity to hypothetical protein An02g12840 - Aspergillus niger	0	0	0	78.2	69.3	41.2	36.3	0
Pc20g06220	similarity to hypothetical protein SPBC173A.10c - Schizosaccharomyces pombe	0	0	0	64.3	66.8	44.7	25.0	1e-49
Pc20g06520	strong similarity to hypothetical protein nc003268.1 - Neurospora crassa	1e-23	AC135027.3	gene: "CU1261C08.3"; product: "SR-rich pre-mRNA splicing a	0	345.4	185.6	94.4	14e-55
Pc20g07120	strong similarity to hypothetical protein An06g00470 - Aspergillus niger	0	0	0	77.8	71.5	37.2	13.8	0
Pc20g07760	strong similarity to hypothetical protein An01g01020 - Aspergillus niger	1e-17	BT007762.12	product: "CG4502.9A"; "CG4502.9A"; Drosophila melanog	20.9	12.1	115.0	60.7	0
Pc20g07870	strong similarity to hypothetical protein contig1487.1.1ta_1500wg - Aspergillus fumigatus	0	0	0	251.0	232.4	108.2	72.1	0
Pc20g08120	similarity to hypothetical protein contig1487.1.1ta_1500wg - Aspergillus fumigatus	0	0	0	53.8	61.6	27.9	12.7	2e-33
Pc20g08670	hypothetical protein	0	0	0	50.9	28.8	18.1	16.4	4e-34
Pc20g08960	similarity to hypothetical protein contig1.145_scaffold_11.1ta_440wg - Aspergillus nidulans	2e-10	NC274.25	gene: "77A.260"; product: "related to secretory pathway pro	119.0	115.4	102.2	47.2	7e-44
Pc20g09010	similarity to hypothetical protein YGR280c - Saccharomyces cerevisiae	9e-14	S64615	hypothetical protein YGR280c - yeast [Saccharomyces cerev	433.9	350.4	179.4	100.3	0
Pc20g09160	strong similarity to hypothetical protein An01g0220 - Aspergillus niger	0	0	0	61.4	62.1	31.0	12.0	0
Pc20g09300	weak similarity to S-layer protein - Clostridium thermocellum	5e-09	AEO17014.3	product: "Cell surface protein"; Bacillus cereus ATCC 1460	166.6	111.5	48.2	51.1	0
Pc20g09710	similarity to hypothetical protein An15g08870 - Aspergillus niger	0	0	0	58.9	46.5	24.5	17.4	0
Pc20g09820	similarity to hypothetical protein An01g07110 - Aspergillus niger [putative pseudogene]	0	0	0	915.2	803.7	329.8	180.7	0
Pc20g09820	hypothetical protein	0	0	0	47.6	21.2	20.0	12.0	0
Pc20g10010	weak similarity to hypothetical protein RV619 - Mycobacterium tuberculosis	3e-07	AP050550.1	gene: "lysS2"; product: "putative lysyl-RNA synthetase"; St	47.8	36.5	34.0	12.0	0
Pc20g10260	strong similarity to hypothetical protein An16g07800 - Aspergillus niger	0	0	0	49.7	44.1	25.0	14.5	2e-69
Pc20g10300	similarity to hypothetical protein contig1.123_scaffold_2.1ta_100wg - Aspergillus nidulans	0	0	0	73.0	28.8	30.7	12.0	3e-60
Pc20g10540	strong similarity to 27 kDa antigen - Parascoccidioides brasiliensis	1e-64	PBU41503.1	product: "27 kDa antigen"; Parascoccidioides brasiliensis 27	75.8	37.5	48.8	29.0	1e-133
Pc20g10590	similarity to hypothetical beta-lactamase XP1621 - Xylella fastidiosa	4e-28	AEO17271.166	product: "conserved hypothetical protein"; Bacillus cereus AT	75.8	37.5	48.8	29.0	1e-133
Pc20g10780	weak similarity to hypothetical protein contig1547.1 - Magnaporthe oryzae	7e-07	AY404974.1	product: "Nucleoside diphosphate kinase 4"; Homo sapiens test	306.1	82.1	131.0	41.1	0
Pc20g11330	strong similarity to hypothetical protein An12g05680 - Aspergillus niger	2e-09	C8212	hypothetical protein [imported] - Arabidopsis thaliana	44.0	22.0	12.0	12.0	0
Pc20g11370	strong similarity to hypothetical protein An08g0980 - Aspergillus niger	0	0	0	64.0	40.2			

Pc20g14620	strong similarity to hypothetical protein An2g04380 - Aspergillus niger	3e-05	NCB1406_25	gene: "B1406.250", product: "hypothetical protein"; Neurosp	161.0	210.0	44.4	48.0	0	
Pc20g14660	strong similarity to hypothetical protein An2g04400 - Aspergillus niger	1e-124	NCB24N1_1_2	gene: "B24N1.020", product: "hypothetical protein"; Neuros	91.4	97.0	30.2	30.9	1e-68	0
Pc20g14860	hypothetical protein	0	0	0	227.8	46.7	68.5	37.0	1e-108	99 UNCLASSIFIED PROTEINS
Pc20g15000	strong similarity to hypothetical protein An2g04750 - Aspergillus niger	3e-04	S67612	hypothetical protein YD076c - yeast (Saccharomyces cerev	66.2	32.9	32.8	21.1	0	0
Pc20g15110	strong similarity to hypothetical protein YHR151c - Saccharomyces cerevisiae	5e-17	S48766	hypothetical protein YHR151c - yeast (Saccharomyces cerev	256.2	216.3	110.3	114.9	0	0
Pc20g15240	similarity to hypothetical protein CAD2106.1 - Neurospora crassa	3e-16	NC25E_17	gene: "SEB1.160", product: "related to APCA protein"; Neuros	105.2	138.8	63.8	61.2	0	0
Pc21g00410	hypothetical protein	0	0	0	75.9	36.2	13.2	17.1	0	0
Pc21g00810	strong similarity to hypothetical protein An7g04840 - Aspergillus niger	0	0	0	148.1	82.8	72.6	48.1	3e-68	03.01.03 DNA synthesis and replication
Pc21g00930	strong similarity to hypothetical protein An2g02240 - Aspergillus niger [putative sequencing error]	0	0	0	36.7	60.4	19.3	12.9	0	0
Pc21g01180	strong similarity to hypothetical protein contig_1_144_scaffold_11.1ta_220cg - Aspergillus nidulans	7e-24	T40058	probable chromatin assembly factor subunit - fission yeast (S	239.0	151.4	69.2	80.1	1e-112	99 UNCLASSIFIED PROTEINS
Pc21g01870	weak similarity to 47 kD subunit of DNA-directed RNA polymerase III Rpo31 - Saccharomyces cerevisiae	6e-05	741150	probable DNA-directed RNA polymerase subunit - fission yeas	114.0	105.6	76.9	47.1	0	0
Pc21g02000	strong similarity to hypothetical membrane protein YOL220w - Saccharomyces cerevisiae	7e-27	S67601	probable membrane protein YOL220w - yeast (Saccharomyces	49.5	67.1	26.1	23.2	0	0
Pc21g02030	strong similarity to hypothetical protein contig3_1_part_1.1ta_2370cg - Aspergillus fumigatus	2e-23	NCB10N12_5	gene: "B10N12.050", product: "putative protein"; Neurospora	48.9	36.8	41.6	20.8	0	0
Pc21g03110	strong similarity to hypothetical protein CAD21186.1 - Neurospora crassa	1e-180	NCB1K11_22	gene: "B1K11.220", product: "hypothetical protein"; Neurosp	63.9	294.1	283.0	93.3	4e-28	02 ENERGY
Pc21g03340	strong similarity to hypothetical protein contig_1_114_scaffold_8.1ta_40cg - Aspergillus nidulans	7e-11	NCB11C21_9	gene: "B11C21.090", product: "hypothetical protein"; Neuros	127.8	127.4	59.1	43.8	0	0
Pc21g03360	strong similarity to hypothetical protein An14g04960 - Aspergillus niger	5e-27	T39803	hypothetical protein SPBC19C2.12 - fission yeast (Schizosac	356.6	444.7	220.4	187.9	0	0
Pc21g03680	weak similarity to hypothetical protein An18g0170 - Aspergillus niger	0	0	0	145.9	152.7	63.3	46.8	0	0
Pc21g04700	weak similarity to hypothetical protein An3g00570 - Aspergillus niger	0	0	0	85.9	60.1	33.3	35.1	0	0
Pc21g04810	similarity to hypothetical protein contig42.1ta_2390cg - Aspergillus fumigatus [truncated ORF]	0	0	0	69.1	131.8	26.4	40.6	4e-34	03.03.01.01.11 mitosis
Pc21g04860	strong similarity to hypothetical protein An12g06000 - Aspergillus niger	0	0	0	128.9	12.8	28.8	40.6	2e-40	01.03.16.01 RNA degradation
Pc21g05000	strong similarity to hypothetical protein An13g02200 - Aspergillus niger	2e-05	AB000516_1	product: "DSF p180"; Homo sapiens mRNA for DSF p180,	1186.4	338.7	814.5	254.2	2e-37	06.07 vesicular transport (Golgi network, etc.)
Pc21g05350	strong similarity to hypothetical RNA-binding protein SPAC1E8.06c - Schizosaccharomyces pombe	5e-35	T37786	probable RNA-binding protein - fission yeast (Schizosacchar	140.3	141.4	131.2	65.9	6e-42	06.07 protein modification
Pc21g05410	strong similarity to hypothetical protein contig_1_22_scaffold_2.1ta_330cg - Aspergillus nidulans	3e-05	AF332335_9	gene: "orf15", product: "Orf15"; Clostridium difficile conjuga	66.1	67.7	34.0	24.5	4e-65	01.05.01 C-compound and carbohydrate utilization
Pc21g05480	strong similarity to zinc metalloprotease like protein An10g0760 - Aspergillus niger	3e-22	T18262	S-layer protein - Clostridium thermocellum	64.6	20.1	25.2	23.9	0	0
Pc21g05780	strong similarity to hypothetical 6-Hydroxy-D-ribose oxidase related protein 13E11.250 - Neurospora crassa	1e-48	T48777	6-HYDROXY-D-NICOTINE OXIDASE related protein [imported	52.1	33.8	35.7	16.9	0	0
Pc21g05820	hypothetical protein	0	0	0	319.4	647.3	148.5	120.7	0	0
Pc21g05930	similarity to hypothetical protein An12g06840 - Aspergillus niger	0	0	0	52.3	97.8	20.2	12.9	0	0
Pc21g05980	strong similarity to hypothetical protein An1g02280 - Aspergillus niger	0	0	0	571.2	652.1	490.0	325.9	0	0
Pc21g06130	strong similarity to hypothetical protein An1g02510 - Aspergillus niger	0	0	0	364.7	290.0	240.1	142.8	0	0
Pc21g06260	similarity to hard surface induced protein 3 chp3 - Gliomastera cingulata	3e-18	AF089807_1	gene: "chp3"; product: "hard surface induced protein 3"; Glo	80.3	24.6	29.7	53.9	1e-47	06.13 proteolytic degradation
Pc21g06270	strong similarity to hypothetical protein contig462_2.1ta_530cg - Aspergillus fumigatus	0	0	0	45.6	12.8	12.0	10.0	0	0
Pc21g06290	strong similarity to hypothetical protein contig5_1_part_1.1ta_2590cg - Aspergillus fumigatus	3e-11	NC03G11_13	gene: "93G11.140", product: "related to serine-related nucle	77.3	47.1	28.1	14.8	3e-35	03.01.05.03 DNA recombination
Pc21g07160	strong similarity to hypothetical protein An16g0205 - Aspergillus nidulans	0	0	0	235.3	298.2	62.9	59.4	0	0
Pc21g07450	strong similarity to hypothetical protein contig_1.1ta_1990cg - Aspergillus fumigatus	7e-38	NCB10N12_7	gene: "B10N12.070", product: "related to rRNA processing	160.4	146.1	128.2	71.7	1e-149	99 UNCLASSIFIED PROTEINS
Pc21g07770	hypothetical protein	0	0	0	62.4	37.0	17.7	12.0	0	0
Pc21g07830	strong similarity to hypothetical protein F28J2.020 - Arabidopsis thaliana	1e-164	AY254382_1	gene: "timpA", product: "TimpA"; Emeritella nidulans TimpA f	226.0	145.0	61.4	137.6	1e-37	99 UNCLASSIFIED PROTEINS
Pc21g07910	hypothetical protein	0	0	0	35.3	14.2	12.0	12.1	0	0
Pc21g07920	strong similarity to hypothetical protein F1086.27 - Arabidopsis thaliana	4e-11	AC133308_8	gene: "OSJNB0807806.07", product: "hypothetical protein";	232.8	130.8	108.9	76.0	0	0
Pc21g07980	strong similarity to hypothetical protein An2g04800 - Aspergillus niger	0	0	0	43.3	20.2	29.7	22.0	0	0
Pc21g08290	strong similarity to hypothetical protein contig49.1ta_100cg - Aspergillus fumigatus	0	0	0	370.3	30.4	12.1	17.9	0	0
Pc21g08530	strong similarity to hypothetical protein contig_1_98_scaffold_6.1ta_760cg - Aspergillus nidulans	0	0	0	51.3	84.2	88.7	22.2	4e-44	01.05 C-compound and carbohydrate metabolism
Pc21g08670	strong similarity to hypothetical protein An14g0360 - Aspergillus niger	0	0	0	132.6	131.7	119.7	47.0	0	0
Pc21g08620	weak similarity to hypothetical human microfilament-associated protein 1 homolog SPAC1782.03 - Schizosaccharomyces pombe	2e-31	NCB11H24_3	gene: "B11H24.030", product: "related to microfilament-associ	44.9	62.3	18.3	19.9	2e-73	04.01.09 other rRNA-transcription activities
Pc21g08860	strong similarity to acvA gene expression regulator Pcr84 like protein An13g01050 - Aspergillus niger	0	0	0	87.3	83.8	62.4	40.1	0	0
Pc21g08900	strong similarity to hypothetical protein YOR294w - Saccharomyces cerevisiae	2e-32	S67198	hypothetical protein YOR294w - yeast (Saccharomyces cere	251.6	221.8	153.1	100.1	0	0
Pc21g08970	strong similarity to hypothetical protein An13g0070 - Aspergillus niger	0	0	0	52.4	30.8	30.7	15.2	2e-59	01.05.01 C-compound and carbohydrate utilization
Pc21g09320	strong similarity to hypothetical protein An13g01120 - Aspergillus niger	1e-88	BX042630_12	gene: "B13D15.120", product: "hypothetical protein"; Neuros	66.0	65.5	38.4	31.5	2e-60	01.01.01.11 biosynthesis of the cysteine-aromatic group
Pc21g09760	hypothetical protein	0	0	0	126.4	12.0	25.3	12.0	1e-48	01.05.01 C-compound and carbohydrate utilization
Pc21g10380	strong similarity to hypothetical protein SPAC19A8.06 - Schizosaccharomyces pombe	2e-05	T37955	hypothetical protein SPAC19A8.06 - fission yeast (Schizos	36.7	36.7	15.2	20.5	0	0
Pc21g10660	strong similarity to hypothetical protein mg06600.1 - Magnaporthe grisea	2e-22	AE015689_5	gene: "SO2474", product: "carboxic anhydrase family protein	345.8	417.4	153.5	203.2	2e-54	10 CELLULAR COMMUNICATION/SIGNAL TRANSDUCTION MECHANISM
Pc21g10980	hypothetical protein	0	0	0	533.7	408.4	266.9	309.9	0	0
Pc21g11210	strong similarity to hypothetical protein contig1403_1.1ta_240cg - Aspergillus fumigatus	2e-08	CEAF3131_1	gene: "unc-89", product: "Uncoordinated protein 89, isolumi	32.5	18.3	16.9	18.9	3e-43	04 TRANSCRIPTION
Pc21g11740	strong similarity to hypothetical protein An12g06000 - Aspergillus niger	0	0	0	67.3	60.8	34.7	15.8	0	0
Pc21g11840	strong similarity to hypothetical protein SOM1 - Neurospora crassa	1e-38	AF265233_1	product: "SOM1 protein"; Neurospora crassa SOM1 protein	271.6	228.7	155.2	111.8	0	0
Pc21g11870	hypothetical protein	0	0	0	57.4	35.8	38.2	24.2	0	0
Pc21g11870	strong similarity to hypothetical protein An12g04000 - Aspergillus niger	0	0	0	23.1	22.5	26.1	12.0	2e-71	01.05 C-compound and carbohydrate metabolism
Pc21g12050	weak similarity to hypothetical protein SPAC23G3.04 - Schizosaccharomyces pombe	5e-20	NCB8P8_19	gene: "B8P8.19", product: "conserved hypothetical protein";	135.3	110.8	93.6	46.4	0	0
Pc21g12080	strong similarity to hypothetical protein contig49.1ta_370cg - Aspergillus fumigatus	2e-13	PFMALP61_235	gene: "MALP61.152", product: "ATP-dependent DEAD box h	88.2	64.7	62.7	37.9	0	01.04 phosphate metabolism
Pc21g12410	hypothetical protein	0	0	0	28.4	12.0	12.0	12.0	7e-44	01.04 phosphate metabolism
Pc21g12730	hypothetical protein	0	0	0	45.1	27.1	30.3	14.5	6e-55	01.04 phosphate metabolism
Pc21g12850	weak similarity to telicuclye binding protein 2 homolog a - Plasmodium falciparum	3e-13	PFMAL13_350	gene: "FPF13.0198", product: "telicuclye binding protein 2 h	86.5	87.1	60.0	21.1	0	0
Pc21g13360	similarity to phosphatidylinositol(3)phosphate binding protein P81 - Saccharomyces cerevisiae	8e-09	1980	gene: "YOR313", product: "YOR313 - yeast (Saccharomyces cere	116.3	84.8	50.0	581.9	1e-35	40.10 nucleus
Pc21g13450	strong similarity to hypothetical protein contig_1_169_scaffold_15.1ta_700cg - Aspergillus nidulans	0	0	0	33.0	12.0	12.0	12.0	0	0
Pc21g13470	weak similarity to hypothetical protein d1042010.5 - Homo sapiens	6e-16	AB037859_1	gene: "K1A1438", product: "K1A1438 protein"; Homo sapie	1846.8	1435.1	922.7	588.2	0	0
Pc21g13480	strong similarity to hypothetical protein An12g01200 - Aspergillus niger	2e-31	CNG507XX_29	data centromeric region sequence from BAC DP15B03, DP3	65.5	62.1	30.2	12.1	1e-154	40 SUBCELLULAR LOCALISATION
Pc21g13910	strong similarity to hypothetical protein contig1483_1.1ta_1660cg - Aspergillus fumigatus	0	0	0	137.5	104.2	47.6	46.5	4e-42	05 PROTEIN SYNTHESIS
Pc21g14400	weak similarity to hypothetical membrane protein YMR260w - Saccharomyces cerevisiae	1e-143	354478	probable membrane protein YMR260w - yeast (Saccharomyce	419.4	296.1	129.8	109.4	0	0
Pc21g14530	weak similarity to hypothetical protein YBL051c - Saccharomyces cerevisiae	1e-78	NCB15B24_8	gene: "B15B24.070", product: "conserved hypothetical protein	99.8	80.8	67.2	37.8	0	0
Pc21g14590	similarity to hypothetical protein contig_1_168_scaffold_14.1ta_210cg - Aspergillus nidulans	0	0	0	40.0	13.4	17.5	12.0	0	0
Pc21g14850	strong similarity to hypothetical protein F3E22.6 - Arabidopsis thaliana	1e-04	AC032912_6	gene: "F3E22.6"; Arabidopsis thaliana chromosome II BAC	603.8	275.3	406.8	179.4	0	0
Pc21g15070	hypothetical protein [putative pseudogen]	0	0	0	32.9	17.8	12.0	13.9	0	0
Pc21g15280	hypothetical protein	0	0	0	221.2	208.2	122.2	71.6	2e-65	01.06 lipid, fatty-acid and isoprenoid metabolism
Pc21g15540	similarity to hypothetical protein CAD11786.1 - Neurospora crassa	2e-10	NC104H10_6	gene: "104H10.050", product: "hypothetical protein"; Neuros	147.8	110.8	85.6	47.7	0	0
Pc21g15470	strong similarity to hypothetical protein contig1480_3.1ta_1000cg - Aspergillus fumigatus	3e-10	AC084196_2	gene: "YS05D8.3", product: "hypothetical protein YS05D8.3"	29.8	68.1	26.4	12.9	0	0
Pc21g16510	similarity to hypothetical protein 1233_scaffold_3.1ta_1120cg - Fusarium graminearum	0	0	0	22.7	25.5	12.0	12.3	4e-38	04 TRANSCRIPTION
Pc21g16750	similarity to hypothetical protein An13g01540 - Aspergillus niger	0	0	0	549.1	367.4	268.4	283.3	0	0
Pc21g16870	strong similarity to hypothetical protein smk1 - Saccharomyces mikatae	1e-27	HHYD08	heat shock protein DDR48 - yeast (Saccharomyces cerevisiae	724.8	1371.7	648.4	501.9	0	0
Pc21g17260	similarity to hypothetical protein An16g1460 - Aspergillus niger	0	0	0	80.8	64.5	38.6	28.7	0	0
Pc21g17270	weak similarity to hypothetical protein contig5_1_part_1.1ta_850cg - Aspergillus fumigatus	1e-14	AB007650_12	Arabidopsis thaliana genomic DNA, chromosome 5, P1 clone	121.7	106.9	65.9	47.3	1e-123	40.02 plasma membrane
Pc21g17480	similarity to hypothetical protein MU422.14 - Arabidopsis thaliana	2e-53	T40083	hypothetical protein SPBC767.09 - fission yeast (Schizosac	287.9	230.3	222.5	146.2	2e-85	05.15 extracellular transport, exocytosis and secretion
Pc21g17570	weak similarity to hypothetical membrane protein YOR154w - Saccharomyces cerevisiae	0.0	NCB7A16_12	gene: "B7A16.120", product: "conserved hypothetical protein"	159.5	150.5	74.5	68.2	2e-73	11.01 stress response
Pc21g17600	strong similarity to hypothetical conserved protein B7A16.130 - Neurospora crassa	0.0	NCB7A16_13	gene: "B7A16.130", product: "related to low-temperature lab	95.8	93.0	104.3	43.7	3e-39	13.07 cell adhesion
Pc21g17610	strong similarity to hypothetical protein B7A16.140 - Neurospora crassa	2e-72	NCB7A16_13	gene: "B7A16.140", product: "related to low-temperature lab	143.3	138.2	129.2	51.9	0	0
Pc21g17760	strong similarity to hypothetical protein An10g0470 - Aspergillus niger	5e-28	NCB10N14_8	gene: "B10N14.080", product: "hypothetical protein"; Neurosp	99.8	66.4	50.0	43.3	1e-68	01.05 C-compound and carbohydrate metabolism
Pc21g17930	strong similarity to hypothetical protein contig1485_2.1ta_210cg - Aspergillus fumigatus	0	0	0	643.1	624.3	436.1	291.1	0.0	04.05.01 mRNA synthesis
Pc21g18100	strong similarity to differentially expressed protein SHy31 - Mus musculus	5e-28	BC061488_1	Mus musculus SH3 domain Y5C like 1, mRNA (cDNA clone	52.0	72.1	38.7	26.8	6e-45	01.03.16.01 RNA degradation
Pc21g18250	similarity to suppressor of 5 creasease virus mutation R91 - Saccharomyces cerevisiae	1e-168	YAHIE_ICHPO	HYPOTHETICAL PROTEIN C2ZF3.14C N CHROMOSOME I	67.8	64.2	38.2	21.8	0	0
Pc21g18310	similarity to hypothetical single-stranded TG-1 binding protein tog - Schizosaccharomyces pombe	7e-75	AF403297_1	product: "glycine-rich protein"; Coccidioides immitis glycine	919.9	1046.0	565.0	330.0	1e-147	06.13 proteolytic degradation
Pc21g18360	hypothetical protein	0	0	0	39.8	34.1	31.0	12.0	0	0
Pc21g19350	similarity to 2MDA.1 protein - Caenorhabditis elegans	2e-38	AY130758_1	gene: "isof", product: "2MDA.1 protein"; Caenorhabditis eleg	234.9	260.1	177.9	111.8	3e-41	01.05 C-compound and carbohydrate metabolism
Pc21g19410	similarity to hypothetical protein contig3_1_part_1.1ta_3240cg - Aspergillus fumigatus	0	0	0	99.5	55.0	44.1	35.1	0	

Pc22g06930	similarity to hypothetical protein contig_1.93_scaffold.6.fts_7700g - Aspergillus nidulans	9e-06	AF11697.26	product: "putative methylase"; Rhodococcus equi virulence p	1140.1	1851.8	509.0	292.8	2e-50	01.20 secondary metabolism	
Pc22g07080	strong similarity to hypothetical protein contig492.0.fts_5610g - Aspergillus fumigatus	0	0	0	161.0	203.8	128.0	74.1	0	0	
Pc22g07170	strong similarity to cDNA O-methyltransferase msl-1 - Aspergillus parasiticus	2e-45	AB022905.1	gene: "msl-1"; product: "O-methyltransferase"; Aspergillus par	58.9	110.6	34.8	32.3	6e-73	06.04 protein targeting, sorting and translocation	0
Pc22g07180	strong similarity to hypothetical protein contig_1.115_scaffold.8.fts_700g - Aspergillus nidulans	9e-07	NC07823.13	gene: "BPM23.130"; product: "related to L-fucose permease";	20.9	51.5	22.3	12.0	0	0	
Pc22g07310	strong similarity to hypothetical protein An02g0130 - Aspergillus fumigatus	4e-08	PF492955.50	gene: "PP02955"; product: "hypothetical protein"; Plasmodi	377.0	860.3	252.3	166.2	0	0	
Pc22g08420	similarity to hypothetical protein contig492.0.fts_5200g - Aspergillus fumigatus	0	0	0	200.6	200.2	134.6	60.9	2e-72	06.04 protein targeting, sorting and translocation	0
Pc22g09310	hypothetical protein	0	0	0	121.6	110.4	98.4	52.8	0	0	
Pc22g09430	strong similarity to hypothetical protein SPAC2323.05c - Schizosaccharomyces pombe	4e-32	T02179	yeast bud pattern determination protein Rax1p homolog [Imp	406.3	398.6	192.4	188.4	0	0	
Pc22g09470	strong similarity to hypothetical protein SPAC334.1.02c - Schizosaccharomyces pombe	5e-75	T36648	hypothetical protein SPAC324.1.02c - fission yeast (Schize	51.9	60.9	52.5	12.0	0	0	
Pc22g09490	strong similarity to hypothetical protein contig469.0.fts_3500g - Aspergillus fumigatus	0	0	0	68.8	75.1	33.8	30.1	0	0	
Pc22g094710	strong similarity to hypothetical protein contig12.fts_2310g - Aspergillus fumigatus	0	0	0	253.1	254.6	174.1	96.4	0	0	
Pc22g09730	strong similarity to hypothetical intracellular protease/amidase related enzyme of the ThuJ family CAC2826 - Clostridium acetobutylicu	1e-35	G07247	gene: "CAC2826"; product: "intracellular protease/amidase related enzyme (ThuJ family) [e	54.5	65.5	27.4	31.4	3e-81	01.01.99 other amino acid metabolism activities	0
Pc22g09910	strong similarity to hypothetical protein An13g01780 - Aspergillus niger [putative pseudogene]	7e-05	AB020515.4	product: "AKG2 protein-like"; Arabidopsis thaliana genomic D	62.1	42.4	56.1	12.0	0	0	
Pc22g09480	strong similarity to hypothetical oxidoreductase PA0147 - Pseudomonas aeruginosa	2e-34	C83828	probable oxidoreductase PA0147 [imported] - Pseudomonas	59.1	47.6	38.3	12.0	3e-65	10 CELLULAR COMMUNICATION/SIGNAL TRANSDUCTION MECHANISM	0
Pc22g09580	similarity to hypothetical protein contig_1.2_scaffold.1.fts_1500g - Aspergillus nidulans [truncated ORF][putative pseudogene]	0	0	0	200.6	42.7	19.7	17.9	0	0	
Pc22g09590	strong similarity to hypothetical protein contig12.fts_1730c - Aspergillus fumigatus	2e-10	CEAF3131.1	gene: "unc-89"; product: "Uncoordinated protein B9, isoform b	1061.9	1103.0	498.8	196.8	0	0	
Pc22g09570	hypothetical protein	0	0	0	136.2	125.5	66.8	53.7	1e-108	99 UNCLASSIFIED PROTEINS	0
Pc22g09800	similarity to hypothetical protein An04g05790 - Aspergillus niger	0	0	0	59.5	55.5	28.2	18.1	3e-40	03.01.09 DNA restriction or modification	0
Pc22g07070	similarity to hypothetical protein B2311.280 - Neurospora crassa	3e-74	T51916	hypothetical protein B2311.280 [imported] - Neurospora cras	56.9	57.8	28.1	22.6	9e-72	03.03 cell cycle	0
Pc22g07200	weak similarity to hypothetical protein YDR389w - Saccharomyces cerevisiae	2e-29	SPAC204H.1	gene: "SPAC204H.01"; "SPAC631.03"; S.pombe chromosom	133.4	105.7	118.4	58.7	1e-154	04 TRANSCRIPTION	0
Pc22g08340	strong similarity to hypothetical protein SPBC2943.09 - Schizosaccharomyces pombe	1e-133	BN048286.16	gene: "B19P44.170"; product: "conserved hypothetical protein	105.7	85.5	80.0	34.9	0	0	
Pc22g08370	strong similarity to PHD-finger protein like protein An02g14310 - Aspergillus niger	4e-63	BT00941.1	gene: "CG1033"; product: "SDO4170p"; Dirosophila melanog	143.9	135.5	69.5	67.8	0	0	
Pc22g08600	strong similarity to hypothetical protein SPOC338.13 - Schizosaccharomyces pombe	2e-58	T41730	hypothetical protein SPOC338.13 - fission yeast (Schizosac	55.1	51.1	31.2	29.0	0	0	
Pc22g08660	hypothetical protein	0	0	0	28.9	88.8	12.0	12.0	0	0	
Pc22g08830	hypothetical protein	0	0	0	61.8	52.7	29.8	31.4	0	0	
Pc22g08850	hypothetical protein	0	0	0	372.6	483.3	204.8	162.4	0	0	
Pc22g08930	strong similarity to hypothetical protein contig2.fts_1040c - Aspergillus fumigatus	0	0	0	49.5	47.9	21.6	12.0	0	0	
Pc22g09000	similarity to hypothetical protein contig52.fts_900c - Aspergillus fumigatus [truncated ORF]	0	0	0	103.7	84.4	54.5	31.2	5e-86	99 UNCLASSIFIED PROTEINS	0
Pc22g09030	hypothetical protein	0	0	0	138.5	120.7	143.0	60.7	0	0	
Pc22g09650	similarity to hypothetical protein YDR489w - Saccharomyces cerevisiae	6e-34	BN048286.10	gene: "B23L4.100"; product: "conserved hypothetical protein	34.9	18.0	18.0	12.0	0	0	
Pc22g10100	hypothetical protein	0	0	0	85.8	84.7	58.2	38.7	1e-50	10.05 transmembrane signal transduction	0
Pc22g10110	strong similarity to hypothetical protein An16g09170 - Aspergillus niger	0	0	0	170.9	150.2	95.6	74.3	0	0	
Pc22g11730	similarity to hypothetical retroelement pol polyprotein - Arabidopsis thaliana	8e-21	F84811	probable retroelement pol polyprotein [imported] - Arabidopsi	27.2	14.7	12.0	12.0	0	0	
Pc22g11800	hypothetical protein	0	0	0	68.4	11.4	30.0	18.0	0	0	
Pc22g11930	similarity to hypothetical protein SPOC483.10c - Schizosaccharomyces pombe	1e-18	T43146	hypothetical protein - fission yeast [Schizosaccharomyces po	191.7	136.7	90.6	60.5	1e-131	99 UNCLASSIFIED PROTEINS	0
Pc22g11950	strong similarity to hypothetical protein contig_1.38_scaffold.2.fts_30w - Aspergillus nidulans	0	0	0	220.8	86.0	103.0	58.2	0	0	
Pc22g12080	strong similarity to hypothetical protein SPBC1822.11c - Schizosaccharomyces pombe	4e-51	T38902	hypothetical protein SPBC1822.11c - fission yeast (Schizos	146.4	194.7	79.6	49.8	0	0	
Pc22g12260	similarity to hypothetical protein contig28.fts_440w - Aspergillus fumigatus	0	0	0	1799.5	1866.3	1195.8	541.0	0	0	
Pc22g12430	strong similarity to hypothetical protein contig_1.30_scaffold.2.fts_190w - Aspergillus nidulans	0	0	0	25.9	36.5	15.5	12.0	3e-29	03.03.01.01.11 mitosis	0
Pc22g12710	weak similarity to hypothetical transcription factor Arg1 - Saccharomyces cerevisiae	3e-06	BN048280.46	gene: "B11E5.480"; product: "putative protein"; Neurospora	91.5	80.7	87.2	38.7	0	0	
Pc22g12740	similarity to hypothetical protein ref - Aspergillus nidulans	2e-53	AF407714.1	gene: "ref"; product: "Rlf6"; Emicella nidulans Rlf6 (rRf)	682.7	721.4	699.8	352.4	0	0	
Pc22g13370	similarity to hypothetical protein SPAC470.13c - Schizosaccharomyces pombe	1e-15	T36817	hypothetical protein SPAC470.13c - fission yeast (Schizosa	117.6	108.9	88.1	42.6	1e-152	99 UNCLASSIFIED PROTEINS	0
Pc22g13450	strong similarity to hypothetical Medusa medA - Aspergillus nidulans	1e-162	AF305599.1	gene: "medA"; product: "Medusa"; Emicella nidulans Med	137.2	144.5	104.5	49.8	0	0	
Pc22g13880	strong similarity to hypothetical protein SPAC182.02c - Schizosaccharomyces pombe	6e-67	NC10410.25	gene: "104H10.25"; product: "conserved hypothetical protein	65.3	81.0	48.6	29.2	0	0	
Pc22g14060	strong similarity to hypothetical protein CAD2120.1 - Neurospora crassa	2e-26	NC068.14	gene: "5E6.160"; product: "conserved hypothetical protein";	167.4	152.2	80.5	84.6	0	0	
Pc22g14380	strong similarity to hypothetical protein An03g0430 - Aspergillus niger	4e-23	NC100H1_16	gene: "100H1.220"; product: "conserved hypothetical protein	22.8	22.8	20.9	20.9	3e-27	05 PROTEIN SYNTHESIS	0
Pc22g14790	hypothetical protein	0	0	0	74.6	57.0	43.0	26.8	0	0	
Pc22g14850	weak similarity to translocation protein SecE7 - Saccharomyces cerevisiae	9e-14	T36335	probable translocation protein - fission yeast (Schizosacchar	149.7	124.7	73.1	69.8	3e-85	25.05.15 myogenesis	0
Pc22g15010	strong similarity to hypothetical protein contig5.fts_1810w - Aspergillus fumigatus	0	0	0	371.3	348.0	154.0	104.0	0	0	
Pc22g15570	strong similarity to hypothetical protein An03g03800 - Aspergillus niger	3e-07	AE003816.4	gene: "shor"; product: "CG1807-PG"; Dirosophila melanog	63.6	67.8	72.1	33.3	3e-52	99 UNCLASSIFIED PROTEINS	0
Pc22g15670	strong similarity to hypothetical protein An03g03930 - Aspergillus niger	0	0	0	221.5	205.0	91.9	53.7	8e-55	11.05.05 violence, disease factors	0
Pc22g15620	strong similarity to hypothetical protein AG036543.1 - Pneumocystis carinii	3e-21	AF30985.5	product: "YH124w-like protein"; Pneumocystis carinii f. sp.	1101.0	819.8	326.5	491.8	0	0	
Pc22g15690	strong similarity to hypothetical protein contig6.fts_1.fts_1570w - Aspergillus fumigatus	2e-20	AY007311.1	product: "cellulase CelA"; Clavibacter michiganensis subsp.	250.2	305.1	71.7	207.3	3e-89	01.05.01 C-compound and carbohydrate utilization	0
Pc22g16040	strong similarity to hypothetical protein An03g04400 - Aspergillus niger	0	0	0	53.0	45.8	23.2	23.8	0	0	
Pc22g16170	weak similarity to hypothetical membrane protein YL151c - Saccharomyces cerevisiae	5e-05	S38174	product: "YL151c"; product: "YL151c"; product: "YL151c";	200.7	244.5	152.6	74.8	0	0	
Pc22g16170	strong similarity to hypothetical protein nc0638.1 - Neurospora crassa	0	0	0	61.4	40.0	17.5	17.7	0	0	
Pc22g16810	strong similarity to hypothetical protein contig5_part.1.fts_1120w - Aspergillus fumigatus	0	0	0	555.1	498.2	391.0	146.8	1e-100	01.06.01 fatty, lipid- and isoprenoid biosynthesis	0
Pc22g16970	weak similarity to hypothetical protein An04g0350 - Aspergillus niger	0	0	0	128.9	116.2	103.3	59.6	0	0	
Pc22g16980	strong similarity to hypothetical long-term assurance protein 1 - Schizosaccharomyces pombe	3e-83	BN048231.30	gene: "B1014.300"; product: "related to protein LACT1"; Neur	331.1	294.1	152.3	153.9	0	0	
Pc22g17510	hypothetical protein	0	0	0	51.2	83.3	30.5	29.5	0	0	
Pc22g18060	similarity to macin like protein An04g01380 - Aspergillus niger	0	0	0	35.9	12.0	12.0	12.0	0	0	
Pc22g18370	strong similarity to hypothetical protein An04g01590 - Aspergillus niger	0	0	0	37.2	18.0	12.0	12.0	2e-25	04.05.01.04 transcriptional control	0
Pc22g18670	similarity to hypothetical zifC3H4C zinc finger protein SPOC548.05c - Schizosaccharomyces pombe	2e-16	SPOC548.5	gene: "SPOC548.05c"; product: "zinc finger protein; zifC3H4C	91.3	77.4	47.4	34.2	0	0	
Pc22g18960	similarity to hypothetical protein SPOC1795.08c - Schizosaccharomyces pombe	4e-21	T41135	hypothetical protein SPOC1795.08c - fission yeast (Schizosa	224.3	184.2	117.4	74.8	0	0	
Pc22g19000	strong similarity to hypothetical protein contig1488.2.fts_470c - Aspergillus fumigatus	0	0	0	36.8	37.3	40.7	18.8	0	0	
Pc22g19140	strong similarity to hypothetical protein An10g0880 - Aspergillus niger	0	0	0	404.0	397.4	200.0	113.6	0	0	
Pc22g19150	strong similarity to hypothetical protein SPBC2582.10 - Schizosaccharomyces pombe	4e-48	T36986	hypothetical protein SPBC2582.10 - fission yeast (Schizosa	302.7	304.1	152.6	161.9	1e-114	03.01.05.01 DNA repair	0
Pc22g19360	strong similarity to hypothetical protein CAD21410.1 - Neurospora crassa	4e-50	NC19F11.6	gene: "19F11.050"; product: "conserved hypothetical protein"	71.5	54.4	40.0	21.5	0	0	
Pc22g19680	strong similarity to hypothetical protein contig31_part.1.fts_3800c - Aspergillus fumigatus	1e-150	BN048229.34	gene: "B20J13.340"; product: "hypothetical protein"; Neuros	655.5	498.5	348.0	128.1	0	0	
Pc22g19870	strong similarity to loaf gene expression regulator like protein An16g0120 - Aspergillus niger	0	0	0	247.4	120.1	16.1	89.0	0	0	
Pc22g20160	weak similarity to hypothetical protein An12g10330 - Aspergillus niger	0	0	0	774.8	380.4	288.8	267.9	5e-60	10 CELLULAR COMMUNICATION/SIGNAL TRANSDUCTION MECHANISM	0
Pc22g20420	weak similarity to exo-alpha-sialidase - Trypanosoma cruzi	4e-04	AF460768.1	product: "glycoprotein precursor"; Cilantro-Congo hemorrhag	289.3	251.7	157.7	111.8	0	0	
Pc22g20450	similarity to hypothetical protein YL020c - Saccharomyces cerevisiae [putative sequencing error]	9e-15	S50791	hypothetical protein YL020c - yeast (Saccharomyces cerev	90.2	70.1	53.5	28.4	0	0	
Pc22g20730	similarity to hypothetical protein CAD3893.1 - Neurospora crassa	8e-66	NC100H1_12	gene: "100H1.150"; product: "hypothetical protein"; Neurosp	80.5	68.0	31.4	51.3	5e-41	99 UNCLASSIFIED PROTEINS	0
Pc22g20740	weak similarity to UDP-N-acetylglucosamine-alpha-1,3-D-mannoside beta-1,4-N-acetylglucosaminyltransferase IV - Homo sapiens	8e-04	AB024911.1	gene: "hGnT-V-H"; product: "UDP-N-acetylglucosamine:alpha	38.9	18.8	12.0	12.0	0	0	
Pc22g20780	weak similarity to hypothetical protein m07603.1 - Arabidopsis thaliana	3e-04	AC039212.6	gene: "F3E22.0"; Arabidopsis thaliana chromosome II BAC	158.5	107.5	50.5	28.9	0	0	
Pc22g20830	similarity to hypothetical protein An02g0610 - Aspergillus niger	0	0	0	48.4	33.5	20.8	12.0	0	0	
Pc22g21110	similarity to hypothetical protein contig31_part.1.fts_2410w - Aspergillus fumigatus	1e-07	AY140094.1	product: "unknown protein"; Arabidopsis thaliana unknown p	217.1	211.9	124.1	87.8	0	0	
Pc22g21420	hypothetical protein	0	0	0	58.9	37.4	41.8	23.9	0	0	
Pc22g21740	hypothetical protein	0	0	0	36.2	40.4	15.9	12.0	3e-90	99 UNCLASSIFIED PROTEINS	0
Pc22g21800	similarity to hypothetical indole-terpene gene cluster protein pavU - Penicillium expansum	4e-18	NC0462_2	gene: "64C2.020"; product: "conserved hypothetical protein";	56.5	72.8	37.0	26.7	0	0	
Pc22g21870	similarity to hypothetical protein contig10.fts_180w - Aspergillus fumigatus	0	0	0	390.7	454.5	237.4	91.1	3e-54	01.05 C-compound and carbohydrate metabolism	0
Pc22g22320	weak similarity to hypothetical protein related to VrkA - Neurospora crassa	0	0	0	183.3	175.4	118.0	86.4	0	0	
Pc22g22490	weak similarity to hypothetical protein contig1497.4.fts_860c - Aspergillus fumigatus	1e-05	AY164486.1	gene: "PK4131"; product: "polycystic kidney disease 1-like 3"	60.1	52.0	36.6	22.8	6e-62	08.07 vesicular transport (Gdg network, etc.)	0
Pc22g23090	hypothetical protein	0	0	0	427.0	252.0	269.2	119.8	0	0	
Pc22g24650	strong similarity to hypothetical protein An15g05550 - Aspergillus niger	0	0	0	83.6	80.4	38.2	12.0	0	0	
Pc22g24660	similarity to hypothetical protein mg07603.1 - Magnaporthe grisea	0	0	0	116.7	111.4	70.7	21.0	0	0	
Pc22g25140	strong similarity to hypothetical protein An02g0610 - Aspergillus niger	0	0	0	525.1	512.0	453.0	234.4	0	0	
Pc22g25480	weak similarity to hypothetical protein contig_1.5_scaffold.1.fts_1230c - Aspergillus nidulans	0	0	0	33.4	12.0	14.1	12.0	6e-87	04.05.01.04 transcriptional control	0
Pc22g25580	strong similarity to hypothetical protein nc06328.1 - Magnaporthe grisea	2e-07	AE								

Pc24g01670	hypothetical protein	0	0	0	236.2	14.3	28.4	177.2	6e-30	01.05.01 C-compound and carbohydrate utilization	0
Pc24g01710	hypothetical protein	0	0	0	87.2	12.3	12.0	125.0	0		
Pc24g01980	similarity to hypothetical protein mg07660.1 - Magnaporthe grisea [putative pseudogene]	4e-07	BC012625_1	product: "Similar to protein phosphatase 1, regulatory (inhibit	104.0	97.6	45.0	37.4	1e-136	04.05.01.04 transcriptional control	
Pc24g02010	hypothetical protein	0	0	0	12.0	12.0	12.0	12.0	1e-125	10.01.01 unspecified signal transduction	
Pc24g02380	similarity to hypothetical protein - Oryza sativa	8e-20	AP003221_23	gene: "P0003009.25", Oryza sativa [aponica cultivar group]	72.9	38.5	57.5	22.9	0		0
Pc24g02510	strong similarity to gag-pol polyprotein - Oryza sativa [putative pseudogene]	7e-35	AC079688_11	gene: "OS.NBa0062M19.11", product: "gag-pol polyprotein"	31.7	27.4	12.9	12.0	0		0
Pc24g02710	hypothetical protein	3e-04	T06613	hypothetical protein REA - Ehrlichia sp. (strain USG3)	598.1	398.7	256.5	143.7	0		0
Pc24g02730	strong similarity to hypothetical protein Ar08g11530 - Aspergillus niger	0	0	0	69.5	12.0	12.0	12.0	0		0
Pc24g02730	strong similarity to hypothetical protein Ar08g11530 - Aspergillus niger	0	0	0	69.5	12.0	12.0	12.0	#N/A		#N/A
Pc24g02760	weak similarity to hypothetical membrane protein YL151c - Saccharomyces cerevisiae [putative pseudogene]	0	0	0	158.5	19.1	20.0	18.9	#N/A		#N/A

Supplementary Table 17. Transcription Factors involved in b-lactam biosynthesis. (a) Pfam trusted matches (domains scoring higher than the gathering threshold)

ORF code	Description of putative <i>P. chrysogenum</i> ORF	Regulation of b-lactam biosynthesis	Transcription Factor	Transcription Factor Domain (start-end) (a)	Binding sites upstream of the pcbAB, pcbC and penDE genes	Average transcript levels @			
						WIS - PAA	WIS + PAA	DS - PAA	DS + PAA
Pc20g13880	strong similarity to catabolite repressor creA - <i>Aspergillus niger</i>	CARBON SOURCE	CreA. Repressor involved in carbon catabolite repression	C2H2 (62-84), (90-114)	SYGGRG binding regions: pcbAB: 5; pcbC: 1; penDE: 7	243.5	195.3	159.9	111.5
Pc20g13890	similarity to hypothetical DNA-binding protein creA – <i>Aspergillus oryzae</i>			No domain identified		72.9	91.9	65.2	64.3
Pc22g17640	weak similarity to catabolite repressor creA - <i>Aspergillus niger</i>		CreC. Repressor involved in carbon catabolite repression	No domain identified		45.4	52.8	73.2	36.7
Pc12g09670	strong similarity to hypothetical protein involved in carbon catabolite repression creC - <i>Aspergillus nidulans</i>			WD40 domain, G-beta repeat (319-357), (361-399), (403-446)		145.2	172.0	125.9	136.3
Pc12g11600	GATA transcription factor nreB - <i>Penicillium chrysogenum</i>	NITROGEN SOURCE	NreB. Regulator of nitrogen metabolite repression	GATA zinc finger (17-51)	GATA binding regions: pcbAB: 3; pcbC: 2; penDE: 6	76.4	78.6	68.1	43.3
Pc22g24480	regulator of nitrogen metabolite repression nre - <i>Penicillium chrysogenum</i> [putative sequencing error]			Nitrogen regulatory protein AreA N terminus (1-73), GATA zinc finger (665-699)		182.5	149.6	144.4	113.1
Pc18g00420	transcription factor pacC - <i>Penicillium chrysogenum</i>	pH	PacC. Mediates regulation of genes in response to ambient pH	C2H2 (94-118), (124-146)	GCCARG binding regions: pcbAB: 3; pcbAB: 3; penDE: 3	211.5	171.5	219.2	151.2
Pc04g00010	strong similarity to developmental regulatory protein brIA - <i>Aspergillus nidulans</i>	DEVELOPMENT	BrIA. Required for activation of development	C2H2 (184-208), (214-239)	MRAGGGR binding regions: pcbAB: 1; pcbC: 1; penDE: 1	0.2	0.8	1.3	0.5
Pc06g00470	strong similarity to developmental regulatory protein brIA - <i>Aspergillus nidulans</i>			C2H2 (321-345), (351-376)		299.2	306.6	69.6	22.2
Pc17g00170	strong similarity to developmental regulatory protein brIA - <i>Aspergillus nidulans</i>			C2H2 (150-174), (180-204)		0.4	0.4	0.6	0.5
Pc22g26080	strong similarity to developmental regulatory protein brIA - <i>Aspergillus nidulans</i>			C2H2 (202-226)		3.6	2.3	4.5	4.5
Pc22g26360	similarity to developmental regulatory protein brIA - <i>Aspergillus nidulans</i> [putative pseudogene]			No domain identified		0.5	1.8	3.4	1.6
Pc23g00400	strong similarity to developmental regulatory protein brIA - <i>Aspergillus nidulans</i>			C2H2 (210-234), (240-264)		0.4	1.5	1.3	0.6
Pc24g00600	strong similarity to developmental regulatory protein brIA - <i>Aspergillus nidulans</i> [putative sequencing error]			C2H2 (210-234), (240-264)		1.7	0.5	0.9	0.5
Pc24g00840	strong similarity to developmental regulatory protein brIA - <i>Aspergillus nidulans</i>			C2H2 (238-262), (268-293)		0.4	1.3	0.8	0.3
Pc24g01490	strong similarity to developmental regulatory protein brIA - <i>Aspergillus nidulans</i>			C2H2 (284-308)		1.5	3.0	1.2	3.4
Pc24g01720	strong similarity to developmental regulatory protein brIA - <i>Aspergillus nidulans</i>			C2H2 (460-484), (490-515)		n.p.	n.p.	n.p.	n.p.
Pc24g02600	strong similarity to developmental regulatory protein brIA - <i>Aspergillus nidulans</i>			C2H2 (210-234), (240-264)		12.4	10.6	12.4	8.1
Pc24g02650	strong similarity to developmental regulatory protein brIA - <i>Aspergillus nidulans</i> [putative pseudogene]			C2H2 (29-53), (59-83)		5.7	5.4	9.3	19.7
Pc16g09610	strong similarity to protein abaA - <i>Aspergillus nidulans</i>		AbaA. Required for activation of development	TEA/ATTS (68-575)	CATTCT binding regions: pcbAB: 1	22.6	8.9	63.5	4.3
Pc12g01590	strong similarity to protein hapB - <i>Aspergillus nidulans</i>	OTHERS	HAPB, HAPE, HAPC. Heterotrimeric CCAAT-binding complex (PENR1-like)	CBF-B/NF-YA subunit B (228-285) CBF-B/NF-Y1 and archaeal histone (85-150), Core histone H2A/H2B/ archaeal histone (161-169)	CCAAT binding regions: pcbAB: 4; pcbC: 2; penDE: 2	167.7	149.5	119.9	86.6
Pc12g04670	strong similarity to HAPE - <i>Aspergillus oryzae</i>			CBF-B/NF-Y and archaeal histone (46-111)		330.5	311.2	310.2	269.8
Pc14g01630	strong similarity to CCAAT-binding protein hapC - <i>Aspergillus oryzae</i>			RFX. Winged helix transcription factor homologue to CPCR1		339.7	388.1	289.1	316.3
Pc20g01690	transcription factor like protein RFX - <i>Penicillium chrysogenum</i>			RFX DNA-binding domain (222-296)		189.0	153.5	95.4	92.5
Pc16g14010	strong similarity to hypothetical methyltransferase AAC34671.1 - <i>Gibberella zeae</i>			LaeA. Regulator of secondary metabolism		365.6	361.9	373.8	395.3
				Methyltransferase domain (193-285)					

@ Values given are the average of three independent experiments
n.p. No probeset present

Supplementary Table 18. PAA upregulated transporters

orf code	Description of putative <i>P. chrysogenum</i> ORF	Best blast homolog to putative <i>P. chrysogenum</i> ORF			Average transcript levels @			
		Species	Gene code	e-value	WIS - PAA	WIS + PAA	DS - PAA	DS + PAA
Pc13g10900	strong similarity to high-affinity nicotinic acid permease Tna1 - <i>Saccharomyces cerevisiae</i>	<i>Aspergillus fumigatus</i>	BX649605_57	1e-180	12	2661	33	2534
Pc16g02690	strong similarity to benomyl methotrexate resistance protein MDR1 - <i>Candida albicans</i>	<i>Aspergillus fumigatus</i>	BX649607_11	1e-78	12	281	12	910
Pc21g01300	strong similarity to membrane protein Tpo2 - <i>Saccharomyces cerevisiae</i>	<i>Acremonium chrysogenum</i>	ACH487683_1	0.0	12	81	12	591
Pc12g13800	strong similarity to allantoin permease Dal5 - <i>Saccharomyces cerevisiae</i>	<i>Schizosaccharomyces pombe</i>	T41604	5e-80	12	483	12	471
Pc13g06330	similarity to hypothetical membrane protein YOL119c - <i>Saccharomyces cerevisiae</i>	<i>Neurospora crassa</i> DNA	NCB23B10_3	3e-51	12	160	26	355
Pc18g01290	strong similarity to dihydroxyphenylglyoxylate transporter Dtr1 - <i>Saccharomyces cerevisiae</i>	<i>Aspergillus terreus</i>	AF141925_12	3e-71	14	89	31	374
Pc12g13630	strong similarity to allantoin permease Dal5 - <i>Saccharomyces cerevisiae</i>	<i>Neurospora crassa</i>	BX842634_3	0.0	12	86	12	141
Pc13g10030	strong similarity to high-affinity nicotinic acid permease Tna1 - <i>Saccharomyces cerevisiae</i>	<i>Neurospora crassa</i>	NC64C2_19	1e-103	23	487	125	1344
Pc22g08750	strong similarity to mitomycin C translocase mct - <i>Streptomyces lavendulae</i>	<i>Neurospora crassa</i>	BX908808_43	1e-44	12	79	12	121
Pc21g14210	similarity to polyamine transport protein Tpo3 - <i>Saccharomyces cerevisiae</i>	<i>Schizosaccharomyces pombe</i>	T41018	2e-23	12	273	12	111
Pc16g09960	strong similarity to hypothetical protein contig_1_43_scaffold_2.tfa_610cg - <i>Aspergillus nidulans</i>	<i>Bdellovibrio bacteriovorus</i>	BX842647_291	6e-48	12	63	12	91
Pc21g19470	strong similarity to mitochondrial succinate-fumarate transporter Sfc1 - <i>Saccharomyces cerevisiae</i>	<i>Neurospora crassa</i>	BX897673_8	1e-125	25	344	48	361
Pc21g09220	strong similarity to fluconazole resistance transporter Flr1 - <i>Saccharomyces cerevisiae</i>	<i>Aspergillus fumigatus</i>	BX649607_11	4e-90	15	402	158	1119
Pc20g14390	strong similarity to mitochondrial phosphate transport protein G7 - <i>Glycine max</i>	<i>Saccharomyces cerevisiae</i>	S50556	1e-97	12	66	36	233
Pc20g06200	strong similarity to hypothetical membrane protein YIL166c - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	S50361	1e-118	12	26	96	526
Pc12g14890	strong similarity to fluconazole resistance protein FLU1 - <i>Candida albicans</i>	<i>Schizosaccharomyces pombe</i>	T41018	1e-123	12	67	18	95
Pc21g12990	strong similarity to polyamine transport protein Tpo1 - <i>Saccharomyces cerevisiae</i>	<i>Neurospora crassa</i>	BX908812_25	1e-149	12	93	40	202
Pc20g00130	strong similarity to myo-inositol transport protein Itr2 - <i>Saccharomyces cerevisiae</i>	<i>Neurospora crassa</i>	NCB17B1_7	1e-141	168	568	150	679
Pc21g05850	strong similarity to allantoin permease Dal5 - <i>Saccharomyces cerevisiae</i>	<i>Schizosaccharomyces pombe</i>	T41604	1e-84	22	101	127	554
Pc16g00500	strong similarity to hypothetical membrane protein YIL166c - <i>Saccharomyces cerevisiae</i>	<i>Neurospora crassa</i>	NC64C2_19	1e-43	33	71	178	717
Pc22g05400	strong similarity to hypothetical monocarboxylate permease Espb6 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	S38065	4e-76	23	89	67	267
Pc20g04510	similarity to multidrug resistance protein Hof1 - <i>Saccharomyces cerevisiae</i>	<i>Schizosaccharomyces pombe</i>	T39346	2e-39	12	44	12	42
Pc12g11990	strong similarity to fluconazole resistance transporter Flr1 - <i>Saccharomyces cerevisiae</i>	<i>Ustilago maydis</i>	UMPLOC_5	3e-79	54	172	63	214
Pc22g14600	strong similarity to ATP-binding cassette multidrug transport protein atrB - <i>Aspergillus nidulans</i>	<i>Botryotinia fuckeliana</i>	AB028872_1	0.0	35	399	176	567
Pc22g03710	strong similarity to hypothetical membrane protein YBR043c - <i>Saccharomyces cerevisiae</i>	<i>Neurospora crassa</i>	NCB8P8_18	1e-168	14	50	26	82
Pc22g20580	strong similarity to multidrug resistance protein Qdr1 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	S49888	9e-67	12	141	53	163
Pc20g08470	strong similarity to protease Mch5 - <i>Saccharomyces cerevisiae</i>	<i>Neurospora crassa</i>	NCB23B10_3	8e-65	21	65	78	239
Pc16g00610	strong similarity to hypothetical membrane protein YMR088c - <i>Saccharomyces cerevisiae</i>	<i>Podospira anserina</i>	CNS07TIX_2	4e-91	55	122	220	657
Pc13g15950	strong similarity to choline permease Hnm1 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	S11175	1e-59	12	62	12	34
Pc21g14260	strong similarity to high-affinity nicotinic acid permease Tna1 - <i>Saccharomyces cerevisiae</i>	<i>Schizosaccharomyces pombe</i>	SPBC1683_12	1e-120	44	88	407	1043
Pc06g01070	strong similarity to membrane protein Tpo2 - <i>Saccharomyces cerevisiae</i>	<i>Neurospora crassa</i>	NCB19C19_12	1e-122	12	58	25	62
Pc18g05780	strong similarity to multidrug resistance protein frn1p - <i>Schizosaccharomyces pombe</i>	<i>Schizosaccharomyces pombe</i>	T40380	9e-79	101	284	219	530
Pc21g12380	strong similarity to hypothetical neutral amino acid permease - <i>Neurospora crassa</i>	<i>Neurospora crassa</i>	S47892	2e-66	33	75	57	134
Pc21g05550	strong similarity to multidrug resistance protein frn1p - <i>Schizosaccharomyces pombe</i>	<i>Neurospora crassa</i>	NCB13O8_14	5e-70	43	156	45	104
Pc16g12280	strong similarity to membrane protein Tpo2 - <i>Saccharomyces cerevisiae</i>	<i>Aspergillus fumigatus</i>	BX649607_11	0.0	38	291	163	364
Pc18g03010	strong similarity to choline permease Hnm1 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	S11175	1e-75	60	125	37	75

@ Values given are the average of three independent experiments

Supplementary Table 19. Upregulated Transporters in DS17690 vs Wisconsin and +PAA vs -PAA

orf code	Description of putative <i>P. chrysogenum</i> ORF	Best blast homolog to putative <i>P. chrysogenum</i> ORF			Average transcript levels @			
		Species	Gene code	e-value	WIS - PAA	WIS + PAA	DS - PAA	DS + PAA
Pc16g02690	strong similarity to benomyl methotrexate resistance protein MDR1 - <i>Candida albicans</i>	<i>Aspergillus fumigatus</i>	BX649607_11	1.00E-78	12	281	12	910
Pc21g01300	strong similarity to membrane protein Tpo2 - <i>Saccharomyces cerevisiae</i>	<i>Acremonium chrysogenum</i>	ACH487683_1	0.0	12	81	12	591
Pc13g06330	similarity to hypothetical membrane protein YOL119c - <i>Saccharomyces cerevisiae</i>	<i>Neurospora crassa</i>	NCB23B10_3	3.00E-51	12	160	26	355
Pc18g01290	strong similarity to dihydroxy transporter Dtr1 - <i>Saccharomyces cerevisiae</i>	<i>Aspergillus terreus</i>	AF141925_12	3.00E-71	14	89	31	374
Pc13g10030	strong similarity to high-affinity nicotinic acid permease Tna1 - <i>Saccharomyces cerevisiae</i>	<i>Neurospora crassa</i>	NC64C2_19	1.00E-103	23	487	125	1344
Pc21g09220	strong similarity to fluconazole resistance transporter Flr1 - <i>Saccharomyces cerevisiae</i>	<i>Aspergillus fumigatus</i>	BX649607_11	4.00E-90	15	402	158	1119
Pc20g14390	strong similarity to mitochondrial phosphate transport protein G7 - <i>Glycine max</i>	<i>Saccharomyces cerevisiae</i>	S50556	1.00E-97	12	66	36	233
Pc20g06200	strong similarity to hypothetical membrane protein YIL166c - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	S50361	1.00E-118	12	26	96	526
Pc21g12990	strong similarity to polyamine transport protein Tpo1 - <i>Saccharomyces cerevisiae</i>	<i>Neurospora crassa</i>	BX908812_25	1.00E-149	12	93	40	202
Pc21g05850	strong similarity to allantoin permease Dal5 - <i>Saccharomyces cerevisiae</i>	<i>Schizosaccharomyces pombe</i>	T41604	1.00E-84	22	101	127	554
Pc16g00500	strong similarity to hypothetical membrane protein YIL166c - <i>Saccharomyces cerevisiae</i>	<i>Neurospora crassa</i>	NC64C2_19	1.00E-43	33	71	178	717
Pc22g05400	strong similarity to hypothetical monocarboxylate permease Esp6 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	S38065	4.00E-76	23	89	67	267
Pc20g08470	strong similarity to protease Mch5 - <i>Saccharomyces cerevisiae</i>	<i>Neurospora crassa</i>	NCB23B10_3	8.00E-65	21	65	78	239
Pc16g00610	strong similarity to hypothetical membrane protein YMR088c - <i>Saccharomyces cerevisiae</i>	<i>Podospora anserina</i>	CNS07TIX_2	4.00E-91	55	122	220	657
Pc21g14260	strong similarity to high-affinity nicotinic acid permease Tna1 - <i>Saccharomyces cerevisiae</i>	<i>Schizosaccharomyces pombe</i>	SPBC1683_12	1.00E-120	44	88	407	1043

@ Values given are the average of three independent experiments

Supplementary Table 20. Regions with amplification or deletion in DS17690 vs. Wisconsin54-1255

The copy numbers and length are estimated. Only the genes with a putative functional annotation are listed per region. For most regions there are probe sets recognizing the both strands of large intergenic regions that confirm the copy number

Cluster #	Copy # estimate		Predicted ORFs spanned		Predicted # ORFs	Estimated length	Remarks
	Wis	DS17690	Start (ORF)	End (ORF)			
1	1	-	Pc09g00050	Pc09g00360	~30	~56kb	Pc0900010 is transposable element Pc09g00240 weak similarity to calcium-independent phospholipase A2 – H.sapiens Not expressed in DS17690
2	2	1	Pc12g16490	Pc12g16580	~10	~24kb	Preceded by ORF "strong sim.to norsolorinic acid ketoreductase nor-1" End is border of contig Pc12g16490 strong similarity to cell polarity protein tea1p – S.pombe Pc12g16500 strong similarity to casein kinase-1 homolog hhp1p – S. pombe Pc12g16510 weak similarity to bZIP transcription factor Yap3 – S.cerevisiae Pc12g16520 strong similarity to phosducin homolog Plp2 – S.cerevisiae Pc12g16530 strong similarity to coatomer gamma subunit 2 copg2 - Homo sapiens Pc12g16540 strong similarity to cytosolic aspartate-tRNA ligase Dps1 – S. cerevisiae Pc12g16550 strong similarity to hypothetical phosphatidyl synthase – S. pombe Pc12g16560 similarity to hypothetical protein YDR306c – S.cerevisiae Pc12g16580 strong similarity to zinc-finger transcription factor Rdr1 – S.cerevisiae (all shared probe sets!) Expression ~2-fold higher in Wis than DS17690
3	1	2	Pc13g04710	Pc13g04910	~21	~77kb	Adjacent to StuA developmental regulator Pc13g04720 weak similarity to vacuolar protein Vac7 – S. cerevisiae Pc13g04880 strong similarity to cytoplasmic ribosomal protein S. cerevisiae Pc13g04890 strong similarity to translational regulator HsGCN1 H. sapiens Pc13g04900 strong similarity to RNA helicase like protein A. niger Expression ~2-fold higher in DS17690 than Wis
4	1	2	Pc16g15290	Pc16g15510	~24	~41kb	preceded by transposal element MARS Pc16g15280 Pc16g15310 strong similarity to ORF1 of transposon Ant1 –A.niger Pc16g15440 weak similarity to ankyrin Ank2 - Homo sapiens Pc16g15470 strong similarity to P type ATPase ENA1 Pc16g15490 strong similarity to delta latroinsectotoxin like protein An08g12230 Expression for several of the ORFs is ~2-fold higher in DS17690 than Wis
5	1	6	Pc21g21280	Pc21g21490	~22	~57kb	Pen amplicon Flanked by transposable elements, 21g21030,-40,-60,21g21240 Pc21g21280 strong similarity to methyl sterol oxidase Erg25 - Saccharomyces cerevisiae Pc21g21370 acyl-coenzyme A:isopenicillin N acyltransferase (acyltransferase) AAT PenDE - Penicillium chrysogenum Pc21g21380 isopenicillin N synthase ips PcbC - Penicillium chrysogenum Pc21g21390 alpha-aminoadipyl-cysteiny-valine synthetase pcbAB acvA - Penicillium chrysogenum [putative sequencing error] Expression of several ORFs is 2-12 fold higher in DS17690 than Wis
6	1	-	Pc23g00800	Pc23g01020	~22		preceded by transposon Pc23g00700 Pc23g00930 with similarity to transposase Not expressed in DS17690

Supplementary Table 21. F Effect of gene silencing on b-lactam productivity

Category	Gene	Description	Pathway	Fold change	Fold change	Relative b-lactam titer	
				Wisconsin54-1255 -> DS17690	without PAA -> with PAA	%	SD
control	none (DS17690)	-	-	-	-	100	± 0.09
control	Pc22g15510	strong similarity to ATP-dependent DNA helicase II subunit Ku70 - Mus musculus	NHEJ	0.9	1.1	99.8	± 0.09
1	Pc18g01330	strong similarity to saccharopine reductase LYS3 - Magnaporthe grisea	Lysine	2.2	1.6	no transformants*	-
1	Pc12g14370	strong similarity to lysine permease Lyp1 - Saccharomyces cerevisiae	Lysine	5.3	2.0	101.2	± 0.01
1	Pc16g10020	sulfate permease SutB - Penicillium chrysogenum	Cysteine	2.5	1.6	64.5	± 0.28
1	Pc22g16570	strong similarity to serine O-acetyltransferase cysA - Aspergillus nidulans	Cysteine	1.7	1.5	64.1	± 0.06
1	Pc22g23110	strong similarity to acetolactate synthase precursor ALS - Schizosaccharomyces pombe	Valine	8.4	1.0	no transformants*	-
2	Pc22g24780	strong similarity to 4-coumarate-CoA ligase 4CL - Arabidopsis thaliana	PAA activation	5.7	13.9	98.1	± 0.01
3	Pc20g02040	strong similarity to methicillin resistance gene HmrA - Staphylococcus aureus	Secretion	25.7	323.5	96.0	± 0.09
4	Pc21g04480	strong similarity to peroxisomal integral membrane protein Per8p - Pichia angusta	Peroxisome biogenesis	1.7	1.2	23.1	± 0.00
5	Pc12g13400	strong similarity to cephalosporin esterase - Rhodosporidium toruloides	b-lactam degradation	0.3	0.7	104.0	± 0.04
	Pc22g13680	strong similarity to hypothetical protein contig1495_1.tfa_1650cg - Aspergillus fumigatus	putative isopenicillinN-CoA epimerase	1.1	1.3	121.9	± 0.18
6	Pc16g01770	strong similarity to phenylacetate hydroxylase pahA - Penicillium chrysogenum	PAA degradation	1.3	46.1	95.3	± 0.00
7	Pc13g13200	strong similarity to dark repressor of conidiation velvet veA - Aspergillus nidulans	Regulation of sec metabolism	0.6	0.9	23.6	± 0.02

* = no transformants obtained; knock-out leads probably to an auxotroph, which is lethal on minimal acetamide selection plates

Supplementary Table 22. Orthologous genes from various fungal genome sequences to perform phylogenetic analysis and build species tree

P. chrysogenum	P. marneffeii	T. stipitatus	N. fischeri	A. clavatus	A. terreus	A. nidulans	A. oryzae	A. niger	G. zeae	A. fumigatus	Protein name
Pc22g25570	PMAA_051070	TSTA_021840	NFIA_021750	ACLA_031360	ATEG_02520	AN0272.3	AO090005000779	An01g05030	XP_380532.1	AFUA_1G02880	Phosphotransferase enzyme family domain protein
Pc13g03220	PMAA_053390	TSTA_019540	NFIA_021230	ACLA_031190	ATEG_02545	AN0259.3	AO090005000758	An01g04710	XP_389338.1	AFUA_1G03420	adenylate kinase Adk2, putative
Pc18g02590	PMAA_055450	TSTA_017480	NFIA_021070	ACLA_031030	ATEG_03104	AN4053.3	AO090009000419	An18g04110	XP_380924.1	AFUA_1G03590	importin 11, putative
Pc16g09250	PMAA_054260	TSTA_018720	NFIA_020210	ACLA_030130	ATEG_03746	AN0432.3	AO090003000873	An01g03570	XP_381102.1	AFUA_1G04540	NADH-cytochrome b5 reductase, putative
Pc21g07860	PMAA_024950	TSTA_033650	NFIA_019390	ACLA_029290	ATEG_03094	AN3895.3	AO090009000410	An18g04200	XP_383842.1	AFUA_1G05360	CAIB/BAIF family enzyme
Pc13g02760	PMAA_019960	TSTA_027970	NFIA_018480	ACLA_028290	ATEG_02990	AN4232.3	AO090001000427	An18g05140	XP_384203.1	AFUA_1G06230	ribosome biogenesis protein Ssf2, putative
Pc18g03350	PMAA_026300	TSTA_072610	NFIA_017990	ACLA_027870	ATEG_02957	AN5711.3	AO090001000465	An18g05750	XP_386525.1	AFUA_1G06690	RLI and DUF367 domain protein
Pc12g15080	PMAA_097790	TSTA_043320	NFIA_013440	ACLA_023490	ATEG_00405	AN1074.3	AO090001000337	An08g04390	XP_388527.1	AFUA_1G12070	glycine cleavage system H protein
Pc13g12860	PMAA_098210	TSTA_043060	NFIA_013260	ACLA_023210	ATEG_00423	AN1066.3	AO090001000304	An08g04880	XP_380839.1	AFUA_1G12250	mitochondrial hypoxia responsive domain protein
Pc13g11340	PMAA_068210	TSTA_088490	NFIA_012800	ACLA_022710	ATEG_00465	AN0999.3	AO090012000638	An08g05530	XP_382099.1	AFUA_1G12760	adenylate kinase-associated protein (cap)
Pc13g11860	PMAA_067400	TSTA_087550	NFIA_012140	ACLA_022060	ATEG_00539	AN0675.3	AO090012000545	An08g06440	XP_390392.1	AFUA_1G13370	afatoxin B1-aldehyde reductase GliO-like, putative
Pc16g14390	PMAA_074230	TSTA_094780	NFIA_010480	ACLA_020420	ATEG_00856	AN0834.3	AO090005001238	An01g13070	XP_386820.1	AFUA_1G14940	protein translocation complex component (Npl1), putative
Pc21g22200	PMAA_066770	TSTA_086870	NFIA_008900	ACLA_019050	ATEG_05152	AN0956.3	AO090005001057	An01g10700	XP_387393.1	AFUA_1G16550	dihydrouridine synthase family protein, putative
Pc21g23610	PMAA_075880	TSTA_096320	NFIA_008390	ACLA_018520	ATEG_05095	AN0632.3	AO090005000976	An01g09860	XP_382709.1	AFUA_1G16990	mRNA splicing factor (Prp18), putative
Pc20g05910	PMAA_021980	TSTA_030260	NFIA_033510	ACLA_094520	ATEG_08153	AN7659.3	AO090701000397	An10g00360	XP_390263.1	AFUA_2G01210	ATP dependent RNA helicase (Dbp5), putative
Pc20g04400	PMAA_087690	TSTA_123080	NFIA_034500	ACLA_092690	ATEG_05703	AN4592.3	AO090011000488	An07g06770	XP_390021.1	AFUA_2G02130	fatty acid desaturase, putative
Pc18g04190	PMAA_055810	TSTA_016580	NFIA_035460	ACLA_091730	ATEG_09581	AN10563.3	AO090120000280	An07g07520	XP_384328.1	AFUA_2G03110	alkaline phosphatase Pho8
Pc18g04290	PMAA_055920	TSTA_016480	NFIA_035530	ACLA_091380	ATEG_09575	AN4510.3	AO090120000268	An07g07610	XP_380992.1	AFUA_2G03170	cyclic nucleotide-binding domain protein
Pc18g06270	PMAA_057540	TSTA_014440	NFIA_080890	ACLA_090430	ATEG_03666	AN5226.3	AO090005001538	An07g08810	XP_389550.1	AFUA_2G04080	GPR/FUN34 family protein
Pc13g04590	PMAA_070510	TSTA_090910	NFIA_083870	ACLA_080050	ATEG_09680	AN6176.3	AO090011000883	An05g00810	XP_380686.1	AFUA_2G08190	tubulin-specific chaperone Rbl2, putative
Pc13g04560	PMAA_070660	TSTA_090960	NFIA_083890	ACLA_080070	ATEG_09663	AN4180.3	AO090011000881	An05g00850	XP_380271.1	AFUA_2G08230	MFS transporter, putative
Pc13g04480	PMAA_070690	TSTA_091000	NFIA_083950	ACLA_080130	ATEG_09667	AN6170.3	AO090011000874	An05g00880	XP_390566.1	AFUA_2G08300	DnaJ domain protein, putative
Pc13g04040	PMAA_070950	TSTA_091380	NFIA_084220	ACLA_080430	ATEG_09696	AN6139.3	AO090011000843	An12g03960	XP_386818.1	AFUA_2G08600	1-acylglycerol-3-phosphate acyltransferase (AtaAp), putative
Pc13g03600	PMAA_071300	TSTA_091820	NFIA_084530	ACLA_080810	ATEG_09879	AN10472.3	AO090011000799	An12g04660	XP_387440.1	AFUA_2G08970	thiamine biosynthetic bifunctional enzyme, putative
Pc22g19950	PMAA_087090	TSTA_122460	NFIA_085350	ACLA_068990	ATEG_04429	AN6014.3	AO090011000642	An16g05150	XP_388719.1	AFUA_2G09910	fatty acid activator Faa4, putative
Pc22g20960	PMAA_047680	TSTA_037480	NFIA_085890	ACLA_069530	ATEG_01520	AN9470.3	AO090011000588	An02g06030	XP_384302.1	AFUA_2G10520	urate oxidase Uaz
Pc22g20850	PMAA_048100	TSTA_038000	NFIA_085960	ACLA_069600	ATEG_01513	AN5971.3	AO090011000578	An02g05880	XP_382117.1	AFUA_2G10600	NADH-ubiquinone oxidoreductase 299 kDa subunit, putative
Pc15g00450	PMAA_048570	TSTA_038630	NFIA_086590	ACLA_070280	ATEG_01445	AN5881.3	AO090026000518	An02g03290	XP_389764.1	AFUA_2G11320	conserved hypothetical protein
Pc15g00160	PMAA_048990	TSTA_038950	NFIA_086780	ACLA_070510	ATEG_01423	AN5861.3	AO090026000492	An02g03570	XP_385846.1	AFUA_2G11540	ketoreductase, putative
Pc20g13910	PMAA_092000	TSTA_049820	NFIA_087050	ACLA_070760	ATEG_01121	AN6200.3	AO090026000462	An02g03860	XP_381729.1	AFUA_2G11810	pre-rRNA processing protein Rrp12, putative
Pc20g14410	PMAA_092250	TSTA_049580	NFIA_087290	ACLA_070990	ATEG_01147	AN10806.3	AO090026000435	An02g04180	XP_386935.1	AFUA_2G12110	YagE family protein
Pc20g14430	PMAA_092270	TSTA_049560	NFIA_087310	ACLA_071010	ATEG_01149	AN6312.3	AO090026000433	An02g04200	XP_387432.1	AFUA_2G12130	conserved hypothetical protein
Pc12g07590	PMAA_092870	TSTA_048790	NFIA_088070	ACLA_071710	ATEG_01247	AN6265.3	AO090026000345	An02g01600	XP_384580.1	AFUA_2G12890	small nucleolar ribonucleoprotein complex subunit, putative
Pc12g07260	PMAA_076170	TSTA_096550	NFIA_088310	ACLA_071950	ATEG_01283	AN6244.3	AO090026000270	An02g01970	XP_391055.1	AFUA_2G13130	3' exonuclease family protein (Rrp42), putative
Pc12g13220	PMAA_022660	TSTA_031180	NFIA_089540	ACLA_073120	ATEG_01049	AN2406.3	AO090023000078	An03g03570	XP_388175.1	AFUA_2G14370	conserved hypothetical protein
Pc14g00060	PMAA_099060	TSTA_042050	NFIA_002920	ACLA_062780	ATEG_09071	AN3776.3	AO090103000095	An13g03680	XP_384132.1	AFUA_3G01840	MFS transporter, putative
Pc20g08410	PMAA_004660	TSTA_103160	NFIA_071470	ACLA_033730	ATEG_08962	AN3432.3	AO090020000042	An11g10890	XP_386235.1	AFUA_3G05740	aldose 1-epimerase, putative
Pc20g03160	PMAA_003380	TSTA_101100	NFIA_070650	ACLA_034560	ATEG_08758	AN4774.3	AO090020000339	An11g09700	XP_389624.1	AFUA_3G06600	siroheme synthase, putative
Pc20g02750	PMAA_002730	TSTA_100180	NFIA_070300	ACLA_034910	ATEG_08726	AN4802.3	AO090020000298	An02g13850	XP_389071.1	AFUA_3G06960	60S ribosomal protein L21, putative
Pc12g15750	PMAA_025630	TSTA_073360	NFIA_067520	ACLA_037580	ATEG_04499	AN5008.3	AO090005001272	An16g03410	XP_390969.1	AFUA_3G09780	conserved hypothetical protein
Pc12g15510	PMAA_025890	TSTA_073120	NFIA_067320	ACLA_037810	ATEG_04520	AN4990.3	AO090003000529	An16g03690	XP_387855.1	AFUA_3G09970	vacuolar iron transporter Ccc1, putative
Pc18g00590	PMAA_023810	TSTA_032240	NFIA_066580	ACLA_038600	ATEG_04600	AN4918.3	AO090003000616	An02g06370	XP_380999.1	AFUA_3G10710	conserved hypothetical protein
Pc18g01280	PMAA_021160	TSTA_029300	NFIA_065580	ACLA_039630	ATEG_04715	AN2877.3	AO090003000735	An02g07570	XP_382947.1	AFUA_3G11750	oxysterol binding protein (Osh5), putative
Pc13g09990	PMAA_015840	TSTA_023780	NFIA_064820	ACLA_040320	ATEG_04144	AN3095.3	AO090005000720	An02g08420	XP_390899.1	AFUA_3G12480	conidiophore development protein HymA
Pc13g14200	PMAA_019550	TSTA_027550	NFIA_064180	ACLA_040880	ATEG_04092	AN3178.3	AO090012000818	An02g09020	XP_381003.1	AFUA_3G13150	deacetylase complex subunit Sds3, putative
Pc20g00270	PMAA_028270	TSTA_070570	NFIA_110210	ACLA_045220	ATEG_06826	AN4303.3	AO090023001003	An04g00140	XP_384747.1	AFUA_4G05940	choline phosphate cytidyltransferase Muq1, putative
Pc22g17740	PMAA_027920	TSTA_070900	NFIA_109510	ACLA_045950	ATEG_05558	AN4382.3	AO090023000914	An04g01100	XP_390498.1	AFUA_4G06710	ATP binding protein
Pc22g18450	PMAA_027040	TSTA_071920	NFIA_108970	ACLA_046570	ATEG_05486	AN4434.3	AO090023000805	An04g01650	XP_381829.1	AFUA_4G07250	37S ribosomal protein Rsm25
Pc13g08120	PMAA_058180	TSTA_013640	NFIA_107810	ACLA_047720	ATEG_05361	AN1721.3	AO090023000724	An04g03130	XP_380452.1	AFUA_4G08350	monosaccharide-P-dolichol utilization protein, putative
Pc18g03540	PMAA_056430	TSTA_015860	NFIA_107080	ACLA_048460	ATEG_05305	AN1639.3	AO090023000652	An04g04040	XP_381261.1	AFUA_4G09090	thioredoxin, putative
Pc22g12070	PMAA_097180	TSTA_043980	NFIA_106100	ACLA_049170	ATEG_00748	AN2056.3	AO090003001239	An04g07080	XP_386925.1	AFUA_4G09950	conserved hypothetical protein
Pc22g12080	PMAA_097090	TSTA_044080	NFIA_106090	ACLA_049180	ATEG_00747	AN2055.3	AO090003001238	An04g07090	XP_386912.1	AFUA_4G09960	conserved hypothetical protein
Pc22g05070	PMAA_041980	TSTA_076270	NFIA_104300	ACLA_050970	ATEG_03872	AN3629.3	AO090003001022	An01g07220	XP_388627.1	AFUA_4G11930	formamidopyrimidine-DNA glycosylase, putative
Pc22g05310	PMAA_042160	TSTA_076450	NFIA_104090	ACLA_051170	ATEG_03851	AN3649.3	AO090003000992	An01g07430	XP_381466.1	AFUA_4G12170	50S ribosomal protein L2
Pc21g14680	PMAA_081490	TSTA_115970	NFIA_040040	ACLA_003190	ATEG_09826	AN8049.3	AO090003001313	An02g11200	XP_387085.1	AFUA_5G02080	NADH-ubiquinone oxidoreductase subunit, putative
Pc21g18850	PMAA_080940	TSTA_115540	NFIA_039360	ACLA_002420	ATEG_09241	AN8119.3	AO090102000391	An02g09940	XP_387546.1	AFUA_5G02740	alpha-1,2-mannosyltransferase (Ktr4), putative
Pc21g18830	PMAA_080920	TSTA_115520	NFIA_039340	ACLA_002400	ATEG_09244	AN8117.3	AO090102000393	An02g09910	XP_387523.1	AFUA_5G02760	fatty acid elongase (Gns1), putative
Pc21g15910	PMAA_082230	TSTA_116920	NFIA_038640	ACLA_001700	ATEG_08067	AN8215.3	AO090102000557	An09g05860	XP_387303.1	AFUA_5G03480	methylene tetrahydrofolate reductase
Pc22g13980	PMAA_083530	TSTA_118420	NFIA_038120	ACLA_001180	ATEG_08011	AN8253.3	AO090102000602	An09g06180	XP_389213.1	AFUA_5G04000	proteasome maturation ans ribosome synthesis protein Nop10, putative
Pc12g05480	PMAA_095350	TSTA_045900	NFIA_037890	ACLA_000950	ATEG_07992	AN8273.3	AO090102000625	An09g06650	XP_380820.1	AFUA_5G04210	ubiquinol-cytochrome C reductase complex core protein 2, putative
Pc21g18000	PMAA_084770	TSTA_119810	NFIA_079930	ACLA_010590	ATEG_09344	AN2238.3	AO090102000219	An17g00770	XP_385309.1	AFUA_5G07340	DnaJ domain protein Psi, putative
Pc22g15750	PMAA_084090	TSTA_119180	NFIA_079330	ACLA_012550	ATEG_08330	AN7736.3	AO090701000709	An03g04080	XP_389021.1	AFUA_5G07960	C2H2 finger and ankyrin domain protein, putative
Pc21g10360	PMAA_073410	TSTA_093810	NFIA_076460	ACLA_013770	ATEG_07632	AN2751.3	AO090010000464	An14g03890	XP_389839.1	AFUA_5G10770	topoisomerase II associated protein (Pat1), putative
Pc21g06050	PMAA_033130	TSTA_064500	NFIA_076130	ACLA_014180	ATEG_07684	AN0191.3	AO090026000732	An10g02210	XP_384450.1	AFUA_5G11130	PAXNEB protein superfamily

Pc21g06450	PMAA_030810	TSTA_067430	NFIA_075690	ACLA_014650	ATEG_02113	AN0138.3	AO090026000670	An18g03180	XP_391038.1	AFUA_5G11580	transcription factor TFIH subunit Tfb4, putative
Pc21g06870	PMAA_031000	TSTA_067230	NFIA_075520	ACLA_014820	ATEG_02094	AN0121.3	AO090120000294	An18g02970	XP_389153.1	AFUA_5G11760	hydroxymethylbilane synthase, putative
Pc21g07110	PMAA_031210	TSTA_067000	NFIA_075300	ACLA_015010	ATEG_02076	AN0105.3	AO090120000318	An18g02370	XP_389830.1	AFUA_5G11985	eukaryotic translation initiation factor eIF1a-like protein, putative
Pc21g07430	PMAA_072220	TSTA_092770	NFIA_074990	ACLA_015280	ATEG_02048	AN0075.3	AO090120000344	An18g02020	XP_387356.1	AFUA_5G12260	disulfide isomerase (TigA), putative
Pc21g04300	PMAA_071960	TSTA_092530	NFIA_074670	ACLA_015590	ATEG_02015	AN0056.3	AO090120000378	An14g06670	XP_382665.1	AFUA_5G12530	arrestin (or S-antigen), N-terminal domain protein
Pc21g03150	PMAA_036920	TSTA_055260	NFIA_073060	ACLA_016910	ATEG_06135	AN6920.3	AO090113000079	An14g05310	XP_391008.1	AFUA_5G13890	SNF7 family protein
Pc22g24340	PMAA_018790	TSTA_026810	NFIA_048320	ACLA_098430	ATEG_07240	AN8676.3	AO090120000095	An12g08730	XP_388872.1	AFUA_6G02110	SRF-type transcription factor (Umc1), putative
Pc22g23790	PMAA_018440	TSTA_026470	NFIA_048700	ACLA_098030	ATEG_07208	AN8704.3	AO090120000131	An12g07830	XP_388852.1	AFUA_6G02440	60S ribosomal protein L24a
Pc20g13010	PMAA_017950	TSTA_025990	NFIA_050480	ACLA_096610	ATEG_07054	AN6614.3	AO090701000148	An15g01510	XP_385325.1	AFUA_6G03950	phospholipid-translocating P-type ATPase, putative
Pc20g12570	PMAA_018110	TSTA_026150	NFIA_050630	ACLA_096430	ATEG_07037	AN6599.3	AO090701000133	An15g01330	XP_387080.1	AFUA_6G04090	DUF28 domain protein
Pc12g04760	PMAA_011510	TSTA_005630	NFIA_051780	ACLA_095330	ATEG_05654	AN6500.3	AO090701000014	An15g00080	XP_382679.1	AFUA_6G05200	60S ribosomal protein L28
Pc22g13280	PMAA_073030	TSTA_093480	NFIA_054060	ACLA_082260	ATEG_05813	AN3923.3	AO090001000497	An11g01610	XP_381370.1	AFUA_6G08420	ubiquitin-protein ligase E3 component (UBR1), putative
Pc12g11370	PMAA_029790	TSTA_068650	NFIA_054280	ACLA_083780	ATEG_06604	AN9512.3	AO090001000524	An11g05700	XP_387040.1	AFUA_6G08630	mitochondrial protein, putative
Pc21g20430	PMAA_035630	TSTA_056610	NFIA_056370	ACLA_084330	ATEG_02371	AN2441.3	AO090023000210	An11g00460	XP_384444.1	AFUA_6G10600	ubiquitin-like activating enzyme (UlaA), putative
Pc22g04060	PMAA_080130	TSTA_114700	NFIA_058290	ACLA_085580	ATEG_03285	AN3737.3	AO090009000186	An06g01480	XP_388025.1	AFUA_6G12330	WD domain protein
Pc22g02350	PMAA_077960	TSTA_112390	NFIA_059110	ACLA_086430	ATEG_03352	AN5493.3	AO090003000443	An08g10300	XP_390707.1	AFUA_6G13190	NupC family nucleoside cotransporter
Pc22g00860	PMAA_062870	TSTA_082450	NFIA_059530	ACLA_086870	ATEG_03406	AN5440.3	AO090103000329	An08g08720	XP_390782.1	AFUA_6G13570	cytochrome c peroxidase, putative
Pc21g12230	PMAA_010440	TSTA_006720	NFIA_114740	ACLA_065510	ATEG_01829	AN9108.3	AO090038000578	An12g00660	XP_389216.1	AFUA_7G02010	indoleamine 2,3-dioxygenase family protein
Pc21g11500	PMAA_011130	TSTA_006070	NFIA_115310	ACLA_064960	ATEG_01886	AN9057.3	AO090311000001	An09g03040	XP_380881.1	AFUA_7G02600	conserved hypothetical protein
Pc21g04880	PMAA_069380	TSTA_089580	NFIA_024940	ACLA_005680	ATEG_04279	AN4259.3	AO090026000816	An13g00430	XP_387298.1	AFUA_7G03980	PCI domain protein
Pc21g04340	PMAA_069250	TSTA_089450	NFIA_025290	ACLA_006020	ATEG_04313	AN5675.3	AO090005000117	An13g00870	XP_385911.1	AFUA_7G04320	UBX domain protein (Ubx5), putative
Pc21g08790	PMAA_088500	TSTA_123800	NFIA_025500	ACLA_006230	ATEG_04342	AN6817.3	AO090005000137	An13g01120	XP_391096.1	AFUA_7G04530	alcohol dehydrogenase, zinc-containing
Pc12g06170	PMAA_091190	TSTA_050820	NFIA_026670	ACLA_007150	ATEG_06387	AN6698.3	AO090005000411	An07g02690	XP_385155.1	AFUA_7G05460	conserved hypothetical protein
Pc16g01080	PMAA_060730	TSTA_010590	NFIA_096460	ACLA_042950	ATEG_10389	AN8881.3	AO090010000768	An03g06860	XP_381208.1	AFUA_8G02760	mitochondrial ornithine carrier protein AmcA/Ort1, putative
Pc21g11120	PMAA_060250	TSTA_011290	NFIA_096990	ACLA_057610	ATEG_00039	AN1442.3	AO090103000044	An16g08830	XP_381391.1	AFUA_8G04260	translocation protein (Sec66), putative
Pc20g09960	PMAA_059540	TSTA_012190	NFIA_097700	ACLA_058300	ATEG_00107	AN1491.3	AO090005000655	An16g07940	XP_382708.1	AFUA_8G04880	COP9 signalosome subunit 1 (CsnA), putative

Title

Molecular analysis of a microbial strain improvement paradigm:

Genome sequencing and analysis of *Penicillium chrysogenum* Wisconsin54-1255

SUPPLEMENTARY DATA

Genome assembly. Approximately 99% of the *P. chrysogenum* genome is represented by the 14 largest scaffolds (greater than 100 kb). Previous studies identified at least four chromosomes in *P. chrysogenum*¹. Most fungi contain from 7 to 20 chromosomes^{2,3}, whereas many closely related *Aspergillus* species contain eight chromosomes⁴⁻⁶.

Preliminary attempts to use PCR amplification of the sequence gaps to elucidate their sequence and to link individual contigs, identified that many of gaps consist of repetitive DNA hampering a solid understanding of these.

Genome sequence and analysis. We have identified a total of 145 nuclear genome encoded tRNA genes, corresponding to 19 amino acids (**Table 1**). The tRNA gene(s) for tryptophan is (are) missing and most likely in gaps in the sequence. The tRNA genes are scattered all over the genome, although sometimes small clusters of 2-3 tRNAs are observed. **Supplementary Tables 23 and 24** present the specificity and codon usage of the identified tRNA genes, which differs from the published usage⁷ (**Supplementary Table 25**). Previously, mycovirus sequences were isolated from the type strain NRRL1951⁸, although there was no evidence for a genomic integration. We have detected no mycovirus sequence in the genome sequence of Wisconsin54-1255. As there were already a dozen mutagenesis treatments needed to obtain the Wisconsin54-1255 strain from NRRL1951 the virus might very well be lost.

Fungal genome comparison. The most conserved functional categories between the different fungi are cell cycle and protein synthesis (**Supplementary Fig. 1**). Compared to all other fungal genomes about 65-94% of these *P. chrysogenum* proteins have an ortholog. *P. chrysogenum* proteins involved in cell rescue, transport and metabolism show a larger number of unique proteins. *P. chrysogenum* proteins that have not been assigned to any functional category due to lack of homologies or functionally described protein domains show the lowest number of orthologs in other fungal genomes.

Supplementary Fig. 2 summarizes the functional classification of *P. chrysogenum* specific ORFs and ORFs with orthologs. Proteins related to TCA cycle and respiration are highly conserved over all fungal genomes (**Supplementary Fig. 3**), including *P. chrysogenum*. In contrast, proteins related to biosynthesis of antibiotics, polyketides, metabolism of aliphatic hydrocarbon compounds, and catabolism of aromatic compounds have generally only few orthologs in other fungal genomes.

Drug and ABC transporters of *P. chrysogenum* show only few orthologs in other fungal genomes, whereas electron/hydrogen carrier proteins, lipid transporter and ion channels are more conserved (**Supplementary Fig. 4**).

Life cycle and sex genes. The comparison of the *P. chrysogenum* and *A. fumigatus* Af293 genomes confirmed the presence of over 40 previously identified euascomycete sex genes (data not shown). The divergence of these genes at the amino acid level is similar to average for these genomes (data not shown). The *P. chrysogenum* mating (MAT) locus contains the MAT1 (alpha-domain) gene, which determines the mating type, confirming the recently reported findings⁹.

Horizontal Gene Transfer (HGT). The origin of the isopenicillinN gene *pcbC*. HGT of *pcbC* from Gram-positive bacteria to fungi has been concluded from sequence similarities with the “evolutionary distance argument” and the “topological argument”. In the evolutionary distance argument¹⁰ the similarity at the DNA- and protein level of 4 isopenicillinN synthase (IPNS) and the highly invariable proteins glyceraldehyde-3-phosphate dehydrogenase, triose phosphate isomerase, Hsp70 and Hsp83 was compared, concluding that HGT from a prokaryotic source to a eukaryotic precursor of *Penicillium*, *Aspergillus* and *Acremonium* took place about 370 Mio years ago. This estimate has been based on the respective DNA-sequences and a nucleotide substitution rate of 10^{-9} nucleotide exchanges per site and year¹¹. A similar conclusion was obtained using 5S rRNA gene sequences as internal control^{12,13}. However, when the related deacetoxycephalosporin C synthetase (DAOCS) and IPNS proteins were taken as orthologs (based on an average amino acid identity of 57%) it was concluded to be ordinary evolution with duplication of genes without HGT¹⁴. However, sequence based trees clearly support HGT for the IPNS-encoding genes from Gram-positives to fungi, followed by adaptation to the fungal background. The branching date between the fungal and prokaryotic IPNS-encoding sequences was estimated to be 852 ± 106 Mio

years based on a maximum likelihood approach¹⁵. Still, some questions remain open. To what fungal ancestor did gene transfer take place? Were there several independent transfers, e.g. to *Aspergillus*/*Penicillium* ancestors and *Acremonium*/*Kallichroma* ancestors? Why are α -amino adipoyl-L-cysteinyl-D-valine synthetase (ACVS) and IPNS of *Penicillium* more similar to *A. oryzae* than to *A. nidulans*? If vertical inheritance would be the case, the *pcbAB*- and *pcbC*-related genes should have been lost from all groups of present organisms except the small number of β -lactam producers.

The origin of the ACV synthetase gene pcbAB. As *pcbAB* and *pcbC* are always linked, a common (bacterial) origin is likely. Neighbour Joining (NJ)-trees based on their amino acid structure are very similar (**Supplementary Fig. 5**). Sequence identities/similarities for different ACVS proteins compared to *P. chrysogenum* are:

	Pchr	Aory	Anid	Acep	Ktet	Nlac	Llys
Pchr	100	79/88	67/80	54/70	53/69	44/61	41/59
Aory		100	66/80	55/70	53/69	53/61	52/60
Anid			100	52/68	51/67	43/60	41/59
Acep				100	63/77	43/60	41/58
Ktet					100	42/59	49/58
Nlac						100	53/67
Llys							100

Pchr = *Penicillium chrysogenum*
 Aory = *Aspergillus oryzae*
 Anid = *Aspergillus nidulans*
 Acep = *Acremonium cephalosporium*
 Ktet = *Kallichroma tethys*
 Nlac = *Nocardia lactamdurans*
 Llys = *Lysobacter lactamgenus*

If non-identical domains are aligned (this means module 1, used for amino adipate, against module 2, for cysteine, or 3, for valine), identities shrink to about 30%. Likewise all other Non-Ribosomal Peptide Synthetase (NRPS) systems regardless of bacterial or fungal origin, have identities below 31%, which seems to be a threshold due to conservation of functional motifs. Thus fungal and bacterial ACVS have at least 41% identity at the amino acid level (being the lowest value of the above comparison). For

IPNS, the identities are significantly higher, presumably due to a higher constraint on functionality:

	Pchr	Aory	Anid	Acep	Ktet	Nlac	Llys
Pchr	100	89/93	80/89	76/87	73/84	57/72	55/71
Aory		100	85/92	77/89	74/85	58/73	56/72
Anid			100	74/85	72/84	60/74	56/73
Acep				100	81/87	57/72	55/69
Ktet					100	57/72	55/71
Nlac						100	60/73
Llys							100

Pchr = *Penicillium chrysogenum*
 Aory = *Aspergillus oryzae*
 Anid = *Aspergillus nidulans*
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 Ktet = *Kallichroma tethys*
 Nlac = *Nocardia lactamdurans*
 Llys = *Lysobacter lactamgenus*

Within certain domain regions of ACVS, similar high conservations can be detected. Closely IPNS-related oxoglutarate-Fe(II) oxygenases have again less than 31% identity (*Stigmatella aurantiaca*), which is very similar comparing ACVS to non-related NRPS.

Does the GC-content of these genes point to their bacterial origin? Although sometimes brought forward, there is no evidence for non-fungal GC-content:

Region	Length (bp)	GC content
Upstream	9271	49%
<i>penDE</i> _exon1	711	56%
<i>penDE</i> _intron1	59	37%
<i>penDE</i> _exon2	162	53%
<i>penDE</i> _intron2	68	51%
<i>penDE</i> _exon3	175	54%
<i>penDE</i> _intron3	64	44%
<i>penDE</i> _exon4	46	47%
Intergenic1	1489	51%
<i>pcbC</i>	1005	57%

Intergenic2	1015	53%
<i>pcbAB</i>	11371	54%
Downstream	14714	47%

For *P. chrysogenum* the overall GC content is 48.9%, 52.8% when only considering the exons, 45.3% for introns, and 44.4% for intergenic regions. Here, it is thus only slightly higher. The *penDE* gene, always considered to be the eukaryotic gene of this biosynthetic cluster, also has a somewhat higher GC-content. However, the GC content of *lys2*¹⁶, considered a typical fungal gene of primary metabolism, is with 54% also above the average. The *pcbAB* gene from *A. nidulans* has a GC content of only 50%, compared to a 53.3% average of coding regions (overall genome is 50.3%). The *pcbAB* gene from *A. oryzae* has a GC content of 52 %, and the *penDE* 54%, while the overall genome value is 48.2%; the exons of an adjacent ankyrin-repeat gene have a GC content of 48% GC. So, there are no clear indications for an increased GC content of the penicillin biosynthetic genes of several fungi.

Why is the penDE gene linked to pcbAB and pcbC? *pcbAB* and *pcbC* have been found linked in all known pro- and eukaryotic systems studied so far and represents a fairly large syntenic region due to the size of the NRPS gene. Strikingly, although the genes are linked, their orientation differs in fungi from bacteria, as fungi have a bi-directional promoter region. The acyl transferase (*penDE*) is only linked in systems producing penicillins, which are restricted so far to the mitosporic Trichocomaceae within the class Eurotiomycetes, including strains of Aspergilli and Penicillia. The gene is not found in bacteria and β -lactam producing ascomycetes of the class Sordariomycetes, with the known examples *Acremonium chrysogenum* and *Kallichroma tethys*. In Sordariomycetes an additional bacterial gene of β -lactam clusters is found (*cefEF*), implying multiple gene transfer events¹⁷. Alternatively all genes could have been transferred in a single event and lost in Eurotiomycetes. IPN epimerase is another gene from the bacterial β -lactam pathway with a possible homolog in the *P. chrysogenum* genome, Pc12g11540 (*cefD* *S. clavuligerus*, see **Supplementary Table 6**), which more likely represents a fungal aminotransferase not related to penicillin with some sequence homology to IPN epimerase. Since *penDE* contains 3 introns a eukaryotic origin has been suggested. Association with the β -lactam cluster is thought to be an example of the selfish cluster hypothesis, providing the advantage of process stabilization¹⁸. At least in

industrial selection programmes this biosynthetic cluster did permit the coordinated amplification of all 3 genes¹⁹.

Blastp analysis of acyl-coenzyme A:Isopenicillin N acyltransferase (AT) recovers a set of early branching bacterial and fungal members, where close homologues represent a subset of the fungal branch, restricted to the ACVS-IPNS associated genes in *A. nidulans*, *A. oryzae*, *A. flavus* and *P. chrysogenum*. Functions of other fungal branches are unknown. Most enzymes, also bacterial ones, have the Gly-Cys cleavage site required for autocatalytic activation. A distance tree clearly supports *penDE* as a eukaryotic/fungal gene. The AT-branch could be considered as recruitment of a hydrolase/transferase to facilitate side chain exchange of IPN in the microbody compartment, which also requires IPN import. Both the *P. chrysogenum* and *A. oryzae* AT have C-terminal microbody targeting sequences (ARL and AKL, respectively), while the *A. nidulans* targeting sequence is less obvious (ANI).

Are there other genes of bacterial origin involved in secondary processes?
Arsenate reductase ArsC. The *A. fumigatus* genome hosts 2 orthologous arsenite resistance clusters combining arsenate reductase, thought to be of actinobacterial origin, an arsenite efflux transporter and an arsenic methyltransferase, both fungal genes, and an arsenic resistance protein, presumably proteobacterial origin²⁰. Only the reductase genes contain no introns. Historically, *Penicillium* has been connected with notorious arsenic fungi in 1892 (a rat exposed to the vapor of fungi grown on arsenic compounds was quickly killed – a sort of gaseous antibiotic), although it is not clear if the respective strain has been *Penicillium* or *Scopulariopsis*²¹. In *P. chrysogenum* an arsenite resistance cluster is located between two retrotransposon homologs (Pc06g02160 and Pc06g02260):

Gene-ID	Protein	Introns	Protein Length	Identities
Pc06g02170	ArsH	1	286	<i>A. terreus</i> 80%, <i>A. fumigatus</i> 80%
Pc06g02180	?	0	236	low sim to fungal proteins (>31%)
Pc06g02190	ArsH-fragment	1	110	<i>A. clavatus</i> 32% (315 AA)
Pc06g02200	regulator?	1	328	>60% larger fungal proteins
Pc06g02210	ArsB	4	366	<i>A. fumigatus</i> 85%
Pc06g02220	ArsC	0	134	<i>A. fumigatus</i> 79%, bacterial 72%
Pc06g02230	transcription factor?	1	108	<i>A. fumigatus</i> 34%

Pc06g02240	transcription factor?	1	102	<i>A. fumigatus</i> 33%, large subunit carbamoyl-P-synthase 30%
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Striking is the very high similarity of ArsC with bacterial reductases (72% identity). Phylogenetic trees reveal close branching of the bacterial and fungal enzymes. This could suggest a more recent transfer than the *pcbAB-pcbC* cluster. Closest homologs are found in *Rhodococcus*, *Brevibacterium*, *Frankia* and *Salinispora*. Recently a transposon containing a set of arsenate-resistance genes has been characterized from *Acidibacillus caldus*²². It is remarkable that *P. chrysogenum* and *Aspergilli* have the bacterial reductases, differing from the yeast reductase Acr2p, which was described as the only known eukaryotic enzyme, besides two types of bacterial reductases using either GSH (also Pc16g02220) or thioredoxin²³.

The resistance protein ArsH has also bacterial homologs, e.g. *Agrobacterium* (57% identity). Aligning the *Pseudomonas* ArsH (ZP_00898054, 233 AA) with the *P. chrysogenum* ArsH (286 AA), there is a core region of 61% identity, while *P. chrysogenum* has extra and deviating terminal regions. The 90 amino acid N-terminus is homologous to all the fungal ArsH proteins. Is this a split gene, partially bacterial, partially fungal, with respect to the one intron in some cases? Does gene adaptation include the gain of additional pieces, like *penDE* in the penicillin cluster, besides codon adjustment?

6-methylsalicylic acid (6-MSA) synthases. 6-MSA and the related orsellinic acid are the precursors for various active metabolites, including patulin (various fungi), chlorotricine (*Streptomyces antibioticus*), neocarzinostatin (*Streptomyces neocarzinostaticus*)²⁴⁻²⁶. While there are no bacterial homologs of the *P. chrysogenum* NRPS genes except *pcbAB*, the two 6-MSA synthase genes (Pc16g00370 and Pc22g08170) do have bacterial homologs. This may seem surprising, as the 6-MSA is considered as a typical fungal multienzyme. But all the sequencing projects revealed hundreds of PKS genes, with the top 20 having a similar size and domain organization and grouping into a fungal and a bacterial subcluster. However, the fungal 6-MSA synthases genes fall into the bacterial class of PKS genes²⁷.

Pc22g08170 (cluster 42 in **Supplementary table 4**) has 82% identity to the 6-MSA synthase from *P. griseofulvum*²⁸. A very similar gene cluster is seen in *A. clavatus* with identities mostly in the 74-82% range. The initial steps in the conversion of 6-MSA

to patulin involve a decarboxylation, two hydroxylation and oxidation steps. The *P. chrysogenum* amidohydrolase and the 2 P450 enzymes are presumably involved in this conversion. A gene encoding an isoeopoxidon dehydrogenase²⁹ cannot be identified in the genome, suggesting that *P. chrysogenum* is not a patulin producer. Also, in non-producing strains of *Byssoschlamys* this enzyme is absent³⁰. Bacterial synthases have 42% identity, suggesting horizontal gene transfer. The amidohydrolase Pc22g08120 is another candidate with bacterial homologs (up to 39% identity).

Pc16g00370 shows only 46% identity to the *P. griseofulum* original identified 6-MSA synthase, but 85% identity to a second MSA-type gene of this strain³¹. Similarities with other fungal and bacterial homologs are comparable to Pc22g08170. The cluster structure suggests the synthesis of a prenylated and hydroxylated 6-MSA derivative (cluster 14 in **Supplementary table 4**).

NRPS enzymes. There are 10 Non-Ribosomal Peptide Synthetases (NRPS) in the *P. chrysogenum* genome (included in the table is one hybrid NRPS-PKS):

	Gene Id	Length (AA)	Domain-architecture	Product
ACV-synthetase Siderophores	Pc21g21390	3790	ATCATCATeTe	penicillin
	Pc16g03850	1900	ATCTC	TAF
Dipeptides	Pc22g20400	2076	ATCTC	TAF
	Pc13g05250	5081	ATCATCTCATCTCT	ferrichrome
	Pc21g01710	2104	ATCAT	?
	Pc21g12630	2268	ATCATC	?
	Pc21g15480	2382	ATCATC	?
PKS-associated (equisetin-type)	Pc16g13930	3962	ks-at-mt-ATd	?
Tetrapeptides	Pc13g14330	5267	CATECATCATCATC	?
	Pc16g04690	6064	ATECAATCATECTCT	?
Hexapeptide	Pc21g10790	7287	ATCATCATECATCATCATC	?

A -adenylate domain, T- carrier domain, C-condensation domain, E- epimerization domain, te-thioesterase domain, ks-ketosynthase domain, at-acyltransferase domain, mt-methyltransferase domain, d-dehydrogenase domain (NAD-binding)

Ferrichrome synthetase The ferrichrome synthetase (Pc13g05250, part of cluster 7 in **Supplementary table 4**) represents a type IV synthetase as in *Neurospora crassa*, *Fusarium graminearum*, *Gibberella zeae* and *Aureobasidium pullulans* forming ferricrocin³². In *P. chrysogenum* the second A-domain contributing Ser is missing the major motif SGTGxPKG and is therefore considered as inactive, and just the first Gly-

domain may be used to include small amino acids. These leads to a product shift from ferricrocin to ferrichrome due to domain inactivation, confirming the observation that *P. chrysogenum* forms ferrichrome³³. The Gly code^{34,35} of the first domain DVFELIMHK is identical to the other type IV synthetases, and also to related enzymes from *S. pombe* and *C. heterostrophus*. The N(5)-acetyl-N(5)-hydroxy-L-ornithine (Aho) code of the third adenylate domain, DVLDIGGIGK, fits to a set of 3 related codes, with position 7 being variable (A, F, G). This particular code is found in the *C. heterostrophus* ferricrocin synthetase and suggests that the second A-domain has been inactivated by some recombination event, whereupon the system has been self-repaired by selection for siderophore production. Some fungi may have two types of ferrichrome-synthetases, ascomycetes (*Chaetomium globosum* and *G. zeae* have type II and IV) as well as basidiomycetes (*Ustilago maydis*, type I and II), but apparently most ascomycetes as *P. chrysogenum* have lost each one of the loci. The divergence of the most prominent types II and IV must have been quite ancient, since similarities are only about 32% identity at the amino acid level. Interestingly, most Aspergilli share the type II synthetase with basidiomycetes, while only *A. clavatus* shares the type IV with *P. chrysogenum* (**Supplementary Fig. 6**). Pc13g05250 is linked to two other known siderophore biosynthesis associated genes: Ornithine N5-oxygenase (Pc13g05260) and SidR DHHC-type Zn-finger protein (Pc13g05270). Ornithine-N5-monooxygenases (OMO) are frequently found next to ferrichrome type or triacetylfusarinineC (TAF)-type (see below) NRPS. At first sight type III/IV/V systems cluster with OMO, as well as type I, but not type II. Type II systems have mostly similar adjacent genes to OMO and Sid2.

Extracellular siderophores TAFs are extracellular siderophores found as cyclic or linear variants. The cyclic and open esters of fusigen, fusarinine B and C have been described for *P. chrysogenum*³⁶. The presence of two related synthetases underlines the importance of iron acquisition. Pc16g03850 (1900 AA) (part of cluster 15 in **Supplementary table 4**): the code DVDHGGAVGK is identical to the *A. terreus*, *N. crassa*, *Coccidioides immitis* and *Magnaporthe grisea* triacetylfusarinine synthetases, as is the general structure. The synthetase is linked to a PKS-cluster and contains two transporters. Pc22g20400 (part of cluster 44 in **Supplementary table 4**), 2076 AA: The well-known A-domain code DVDGGGGIGK matches the TAF synthetases from *A. niger* (An03g03520), *A. nidulans* (2086 AA), *A. oryzae* (1932 AA), *A. fumigatus* (2047 and 2083 AA, largely identical), *Botryotinia fuckeliana* (2058 AA) and *Chaetomium globosum*

(1733 AA). The putative gene cluster contains an acetylase and a highly conserved transporter.

Dipeptide synthetases Pc21g01710 (part of cluster 28 in **Supplementary table 4**), similar to An04g06260 of *A. niger* (2196 AA) matching a 2323 AA sequence, with N- and C-terminal deviations and a 58 AA gap. Both specificity regions are similar to *A. fumigatus* enzymes with 2336 and 2210 AA, respectively. The smaller *A. fumigatus* NRPS (2210 AA) has identical codes SARGTVSQLK and DVYFTGGVLK. While the *A. fumigatus* NRPS have the ATCATC structure, no terminal C-domains are detected in the NRPS of *A. niger* and *Penicillium*. Both *A. fumigatus* and two related *A. terreus* NRPS (2610 AA, codes SARDTAAQVK, DAFMLCGILK and 2923 AA, codes SARGTVTQLK, DAQIIGVMVK with the structures CATCATC) has been classified by as “putative unknown siderophore synthetases”, based on a phylogenetic grouping of a set of Aspergilli-NRPS³⁷. Pc21g12630 (part of cluster 35 in **Supplementary table 4**), 2382 AA, represents a cyclodipeptide synthetase, the structure has no significant homologs. The codes DVRSVGAGIK and DIGLGAMVIK have no matches with known NRPS, the second A domain specificity region is related to Ala/Gly/Pro-substrates. Pc21g15480 (part of cluster 37 in **Supplementary table 4**), 2372 AA, represents a cyclodipeptide synthetase, the structure has no significant homologs. The first A-domain code DVRSVGAGIK is identical to Pc21g12630, while the second domain code DSLELVAVVK differs. The linked dimethylallyltransferase and methyltransferase suggest a dimethylallyl-modified methylated cyclodipeptide, or piperazinedione, or diketopiperazine. Roquefortine- and meleagrin-like compounds are candidates, but both are not methylated. Such peptides interfere with bacterial quorum sensing, and could suppress bacterial interactions^{**}. Similarly, diketopiperazines were found to be capable of activating or antagonizing other LuxR-based quorum-sensing systems, such as the N-butanoylhomoserine lactone-dependent swarming motility of *Serratia liquefaciens*. Although the physiological role of these diketopiperazines has yet to be established, their activity suggests the existence of cross talk among bacterial signalling systems. Interestingly, Pc21g12630 is 40-fold overexpressed in the high-producing strain, as well as other members of this cluster. So, together with penicillin production this very well could be a concerted antibacterial response.

Tetrapeptide synthetase Pc13g14330 (part of cluster 10 in **Supplementary table 4**), 5267 AA, represents a cyclotetrapeptide synthetase, the first module epimerises its amino acid. The N-terminal condensation domain could indicate a precursor transfer,

common in bacterial systems, if an associated system providing a carrier protein linked intermediate is available, or this region may just be considered as a remnant from domain deletion events. The codes DSICVVAAVK, DAVLVGAVIK, DVLIMITVVK, DVMFGQAVIK have no match and specificity region analysis provides no clear result, except for module 2 hinting at the Leu/Phe type. Orthologs with 52 and 47% identity are found in *A. terreus* (EAU30156) and *A. clavatus* (EAW15286). All 3 NRPS are linked with transporters, indicating export of the product. No other adjacent genes are shared. The adjacent Pc13g14340, an unknown protein of more than 2000 AA, may be associated with signalling, as it is related to β -transducin regions and NACHT-protein sequences involved in secretion processes. Similar sized unknown proteins can be found in *A. terreus* and *A. clavatus* in different genomic regions. Likewise very often NRPS genes are found next to genes containing ankyrin repeats.

Pentapeptide/hexapeptide synthetases Pc16g04690 (part of cluster 16 in **Supplementary table 4**), 6064 AA, is an ortholog of *A. niger* An08g02310 (54% identity), which has been connected to malformin due to its epimerisation domains and its high similarity to a set of 5 module NRPS resembling the malformin NRPS. A malformin-like product has been reported for *P. roquefortii*³⁸. Similar NRPS are also found in *A. fumigatus* (6229 AA), *A. oryzae* (5199 AA, with the second A-domain deleted), *A. nidulans* (even two with 5935 and 6077 AA) and *A. terreus* (5842 AA). More than 50% identity links this group to a set of 5-module NRPS of the structure ATEC-ATC-ATEC-ATC-ATEC-TCT. Pc21g10790 (part of cluster 33 in **Supplementary table 4**), 7287 AA, shows a 57% identity to a 6885 AA NRPS from *A. oryzae*. This NRPS has an identical structure ATCATCATECTCATCATC, except for loss of the fourth A-domain, suggesting an inactive domain in the *Penicillium* enzyme. The codes of modules 1-3 are identical in *Penicillium* and *A. oryzae*: DVLFAGGVAK, DASFIGVIYK, DVG FVGSIWK, the 5th module is slightly altered DVDEVSSVCK/DVEEASTVSK (*Aory*/*Pchry*, respectively), as is the 6th module in a single position DVACVSAVWK/DIACVSAVWK. All codes are unknown. Despite the inactive A-domain the product could be a cyclohexapeptide, as carrier and condensation domains are present, and *in trans* charging could occur, so that even two adjacent D-amino acids are possible. Alternatively, the product is a cyclopentapeptide with one D-amino acid. An identical product should be present in *A. oryzae*. Since a fatty acid synthase, hydroxylase and aminotransferase are linked, an aliphatic β -amino acid may be involved. Pc16g13930 (part of cluster 20 in **Supplementary table 4**), 3962 AA, is a mixed type PKS-NRPS.

Similar structures are equisetin synthetase (*Fusarium heterosporum*, 3953 AA), a 4034 AA PKS from *Magnaporthe grisea*, an 3821 AA PKS from *Coccidioides immitis*, a 3946 AA PKS from *A. oryzae*, and a 3930 PKS from *A. nidulans*. The A-domain specificity is unknown, the code DIALYGAIK has no match; the specificity region has a weak Arg connection.

PKS enzymes Known polyketide metabolites from *P. chrysogenum* are secalonic acids (octaketides), sorbicillins, sorrentanones (hexaketides), emodic acids (octaketides), chrysogenin (structure not fully elucidated), and the possible PKSIII derived xanthocillins³⁹. Twenty-four putative PKS encoding genes can be extracted from the genome (**Supplementary table 3**). Pc21g16000 can be assigned by similarity to the *A. nidulans* wA and the *A. fumigatus* PksP to conidial yellow pigment biosynthesis (naphthopyrone), and could be the chrysogenin synthase. Pc22g08170 has almost 90% identity to the original 6-MSA sequence from *P. griseofulvum*, and is thus involved in formation of a patulin-like product. The only chalcone synthase type Pc22g09640 could be tentatively related to xanthocillins, also produced by *A. candidus*, *A. chevalieri*, *E. chevalieri*. None of the remaining 21 PKS can be clearly correlated with a product. Pc16g03800 is associated with one of the TAF clusters, and could be involved in synthesis of cis-5-hydroxy-3-methylpent-2-enoic acid, one constituent of fusarinine, the substrate of TAF synthetase.

Amplified chromosomal region with penicillin biosynthetic genes Industrial *Penicillium* strains have a 60-100 kb chromosomal region amplified several times^{19,40}. Sequences and lengths reported¹⁹ were used to extract the exact sequence from the genome. A region of well over 120 kb, covering all reported amplified fragments, is represented in **Supplementary Fig. 7**. Forty-two putative ORFs were identified in the amplified part (4 surrounding putative ORFs are also shown). Twenty-four of these reside in the 57 kb region recently reported in more detail^{41,42}, which is amplified in all industrial strains. This region contains the three genes encoding the key steps in penicillin biosynthesis (ACVS, IPNS and AT). From annotation alone none of the other ORFs can be directly related to penicillin biosynthesis (**Supplementary Table 5**), confirming the recent publications^{41,42}. Some of these ORFs are expressed significantly higher (e.g. more than 2-fold) during penicillinG producing conditions, but this is mostly limited to the Wisconsin54-1255 strain (**Supplementary Table 6** and **Supplementary**

Fig. 8). While the high producing strain DS17690 has 5-7 copies of this region (M.A. van den Berg, unpublished results) it is obvious that the increased gene dosage is not balanced throughout the region. For example ORFs Pc21g21460 through Pc21g21490 show 4- to 8-fold higher mRNA levels, while Pc21g21290 and Pc21g21440 both come close to 30-fold increased levels. As the latter two encode hypothetical proteins this complex regulation effect needs further attention.

The other part of the region is also amplified in strains from the Panlabs lineage^{19,43}. Here, several ORFs with membrane spanning regions are present, including a MFS type transporter, Pc21g21510, homologous to the cercosporin transporter of *Cercospora kikuchii*, which is involved in the secretion and resistance against this PKS-derived toxin⁴⁴. However, as most of the genes in this region it does not show a high transcription level or an increased transcription during penicillin biosynthesis, the exact function remains unclear.

Transcription factors The transcription profiles of the 612 ORFs encoding the putative transcriptional factors showed that approximately half of these are not very actively transcribed (e.g. an average signal <50; **Supplementary Table 26**). Down-regulation under penicillin producing conditions was observed in ORFs putatively encoding important transcription factors such as *abaA*, involved in growth and development; *afIR* (only down-regulated in the DS17690 strain), a transcription factor required for the expression of the aflatoxin pathway genes which negatively regulates the expression of LaeA (a transcription factor directly involved in regulation of secondary metabolism in *Aspergillus* spp.); *amdX* (only down-regulated in the DS17690 strain), a transcription factor regulating the acetamidase-encoding *amdS* gene and *brlA*, involved in growth and development. Up regulated ORFs during penicillin production are *alcR* (only up regulated in the DS17690 strain), a positive regulatory protein for the ethanol regulon, *alcA* and *aldA*; *amdA* (only up regulated in the Wis54-1255 strain) controlling the expression of the acetamidase-encoding *amdS* and *aciA* genes, which allows the utilization of certain amides as carbon and/or nitrogen sources; *arg81*, which mediates the repression of specific arginine biosynthetic genes (ARG) and also functions as an inducer of the arginine catabolic genes (CAR) in response to exogenous arginine; *aro80*, a transcription activator required for the expression of genes involved in the catabolism of aromatic amino acids; a pathway-specific regulatory protein of nitrate assimilation (only up regulated in the DS17690 strain) and SPBC530.05, a transcriptional regulator

similar to the *A. nidulans* *facB* DNA binding protein, which is required for growth on acetate. The strongest responses were observed for uncharacterized factors like Pc22g16820 (e.g. weak similarity to hypothetical transcription regulator SPBC530.05 – *S. pombe*) with a 10-fold signal increase from 155 to 1502 and Pc21g23810 (e.g. strong similarity to hypothetical transcriptional regulator CAF32162.1 – *A. fumigatus*) with a 50-fold signal increase from 13 to 648.

Among the transcription factors reported to control penicillin biosynthesis (like *pacC*, *creA*, *cpcR1*, *laeA*) none were found to be significantly changed in the data analysed. For most of these the *P. chrysogenum* homologues can be readily identified (**Supplementary table 17**). For *creA*, exerting glucose repression, three homologous ORFs can be identified, but Pc20g13880 seems to be the best candidate, which shows a negative mRNA level with increasing penicillin productivity. Although, in all *A. nidulans* *creA* mutants tested, glucose still represses the *ipnA* (=pcbC) gene expression^{45,46}, recent studies in *P. chrysogenum* with antisense *creA* RNA and directed mutation of the CreA binding sites indicate that CreA has an important role in glucose regulation of penicillin biosynthesis (C. Cepeda, F. Fierro and J.F. Martín, unpublished results; L. Cova and M.A. van den Berg, unpublished results). Comparing the transcriptional map of the (extended) amplified region containing the penicillin cluster (**Supplementary Fig. 8**) with the extracted binding sites for transcription factors (**Supplementary Fig. 9**) does not give any further clues for the specific regulation or roles of the co-amplified genes.

Biosynthesis of secondary metabolism is quite often highly regulated. In several examples it was shown that pathway specific transcription factor genes can be physically associated with the biosynthetic genes⁴⁷.

C2H2/Zn(II)2Cys6 transcription factor genes (CMR1-type). Seven putative CMR1-type transcription factors were identified in the genome (**Supplementary table 26**). Each ORF contains two C2H2 DNA-binding domains and one Zn(II)2Cys6 DNA-binding domain. The Zn(II)2Cys6 motif is present at the N-terminal region of CMR1 and the two C2H2 zinc fingers are present N-terminal to the Zn(II)2Cys6 DNA-binding motif corroborating earlier reports⁴⁸.

	C2H2 (Positions)	Zn(II)2Cys6 (Positions)
Pc6g01890	(23-47), (53-76)	(87-124)
Pc12g09540	(17-39), (45-67)	(88-125)
Pc16g05230	(14-36), (42-64)	(78-115)

Pc20g05960	(18-40), (46-68)	(80-117)
Pc21g07180	(37-56), (65-87)	(107-144)
Pc21g07310	(4-28), (34-57)	(69-106)
Pc22g15230	(2-24), (30-52)	(71-109)

Regulators of the P. chrysogenum putative secondary metabolite clusters. From the 47 clusters containing PKS and NRPS only ten contain regulatory genes associated to them. However, all of them remain to been studied at the functional level in *Penicillium chrysogenum*.

Hybrid cluster 11 (Supplementary table 4)

In the hybrid cluster 11 there is one ORF (Pc14g00020) encoding a protein with weak similarity to hypothetical transcription regulator protein from *S. Pombe*.

PKS-NRPS cluster 15 (Supplementary table 4)

In the PKS-NRPS cluster 15 there is one ORF (Pc16g03740) encoding a protein with weak similarity to siderophore biosynthesis repressor sREA from *A. nidulans*.

PKS cluster 17 (Supplementary table 4)

In the PKS cluster 17 there is one ORF (Pc16g04880) encoding a protein wiith weak similarity to hypothetical transcription activator SPAC139.03 from *S. pombe*.

NRPS-like cluster 21 (Supplementary table 4)

In the NRPS-like cluster 21 there is one ORF (Pc18g00330) encoding a protein with strong similarity to hypothetical phd finger transcription regulator from *S. pombe*, one ORF (Pc18g00400) encoding a protein with similarity to hypothetical transcription regulator SPBC530.05 from *S. pombe* and the ORF Pc18g00420, encoding the transcription factor pacC.

PKS cluster 27 (Supplementary table 4)

In the PKS cluster 27 there is one ORF (Pc21g00920) encoding a protein with similarity to transcription activator of lysine pathway Lys14 from *S. cerevisiae*.

PKS cluster 29 (Supplementary table 4)

In the PKS cluster 29 there is one ORF (Pc21g03950) encoding a protein with similarity to hypothetical transcription regulator SPBC530.05 from *S. pombe*.

PKS cluster 31 (Supplementary table 4)

In the PKS cluster 31 there is one ORF (Pc21g04750) encoding a protein with similarity to aflatoxin biosynthesis regulatory protein aflR from *A. parasiticus*.

PKS cluster 32 (Supplementary table 4)

In the PKS cluster 32 there is one OFR (Pc21g05050) encoding a protein with weak similarity to positive regulator qa-1F from *N. crassa* and one ORF (Pc21g05090) encoding a protein with weak similarity to positive regulator of purine utilisation uaY from *A. nidulans*.

PKS cluster 34 (Supplementary table 4)

In the PKS cluster 34 there is one ORF (Pc21g12340) encoding a protein with similarity to hypothetical transcription regulator SPAC139.03 from *S. pombe* and one ORF (Pc21g12360) encoding a protein with strong similarity to hypothetical transcriptional regulator CAF32162.1 from *A. fumigatus*

PKS cluster 42 (6-MSA synthases cluster, Supplementary table 4):

6- methylsalicylic acid (6-MSA) synthases. 6-MSA and the related orsellinic acid are the precursors for various active metabolites, including patulin (various fungi), chlorotricine (*Streptomyces antibioticus*), neocarzinostatin (*S. neocarzinostaticus*).

In the cluster there is one ORF (Pc22g08140) encoding a protein with weak similarity to hypothetical transcription regulator SPBC530.05 from *S. pombe*.

Transporters related to secondary metabolite genes Based on common regulation patterns observed via the MicroArray studies, clusters around several NRPS genes can be identified. Pc21g12630 is predicted to be a cyclodipeptide synthetase, but in the vicinity there are no transporter genes. NRPS Pc21g15480 is an NRPS that is down

regulated under penicillinG producing conditions. The neighboring gene, Pc21g15420, shows strong similarity to the cercosporin transporter CFP of *Cercospora kikuchii*. This gene is regulated in concert with the NRPS and thus might be involved in secretion of the peptide. Pc21g21390 (*pcbAB*; ACV synthetase) is part of the penicillin cluster of three genes (*pcbAB*, *pcbC*, and *penDE*). In contrast to the penicillin cluster in *A. chrysogenum*, the minimal penicillin cluster (e.g. the core 17 kb region covering the three biosynthetic genes) in *P. chrysogenum* does not contain any transporter gene in its proximity. Three transporter genes localize downstream: e.g., Pc21g21510 which is not expressed and shows strong similarity to cercosporin transporter CFP of *C. kikuchii*, Pc21g21530, also not expressed and with strong similarity to allantoin permease Dal5 of *S. cerevisiae*; and Pc21g21590, homologous to the glucose transporter Rco-3 of *N. crassa*, which is expressed at low levels but appears to be down regulated under penicillinG producing conditions (see also **Supplementary table 5**). The latter protein is homologous to the glucose transporter Rco-3 of *N. crassa*.

Two physically close transporter genes are slightly co-upregulated under penicillin producing conditions: Pc22g20390 with a strong similarity to multidrug resistance (MDR) protein AtrD of *A. nidulans* and Pc22g20360 with a strong similarity to siderophore-iron transporter for enterobactin Enb1 of *S. cerevisiae*. In the DS17690 strain Pc22g20390 is up-regulated 3-fold. Recent data shows that it localizes to the plasma membrane as verified by a GFP fusion protein (A. Kovalchuk, unpublished results), and thus seems an interesting candidate for secretion system for penicillin. It should however be stressed that penicillin secretion might be the result of the activity of multitude of MDR-like transporters as the genomic analysis suggests a high level of redundancy in the distribution of such transporters. Most of the common up-regulated genes seem to respond to the high concentration of phenylacetic acid (PAA) used in the fermentation broth. PAA enters the cell mostly by means of passive diffusion, but it is also metabolised⁴⁹ and the conversion products might be secreted. Up to 143 transporter genes are significantly up-regulated in the DS17690 strain versus the Wisconsin54-1255 strain when grown in the presence of PAA. Of this group of transporters, 15 transporter genes seems to be expressed at a higher level in the DS17690 versus Wisconsin54-1255 strain fermentations in the absence of PAA (**Supplementary Fig. 10**). Except for Pc20g14390 that shows homology to mitochondrial phosphate transporters, all other of these 15 up-regulated transporter genes belong to the major facilitator superfamily (MFS) type. These transporters could

be involved in PAA uptake and secretion, precursor uptake and secretion into and out of the microbody and penicillin secretion, or reflect some stress response to the fermentation conditions.

Five (Pc13g10900, Pc16g02690, Pc12g13800, Pc13g06330 and Pc13g10030) of the 36 genes up-regulated under penicillin producing conditions are at least up-regulated 10-fold (**Supplementary table 18**). These likely respond to the added PAA. Pc13g10900 (220- and 77-fold up in the Wisconsin54-1255 and DS17690, respectively) and Pc13g10300 (21- and 11-fold up in the Wisconsin54-1255 and DS17690, respectively) show a strong similarity to high-affinity nicotinic acid permease Tna1 of *S. cerevisiae*. Nicotinic acid and PAA are both small molecules with aromatic rings, suggesting that these two transporters are likely candidates for PAA uptake systems.

Microbodies *Development* All genes encoding proteins essential for microbody formation⁵⁰ (designated peroxins) can be identified in the *P. chrysogenum* genome suggesting that these organelles are formed via conserved machineries. Additionally, all components essential for microbody fission (e.g. Vps1, Dnm1, Fis1^{51,52}) are present, indicating that organelle fission as a mode for microbody multiplication may normally occur.

Stability. Yeast microbody-borne proteins as well as the microbodies themselves have a limited life span and are subject to gradual continuous degradation. In this process both microbody-borne hydrolases (e.g. Lon protease)⁵³ and selective autophagy machineries (pexophagy⁵⁴) play a distinct role. Such degrading AuTophagy (ATG) genes are conserved in the *P. chrysogenum* genome⁵⁵ and shown to play a role in selective microbody turnover (W.H.Meijer *et al.*, unpublished data). A Lon protease with a putative PTS1 is also present in *P. chrysogenum* (Pc21g06860, see **Supplementary table 7**).

Penicillin biosynthesis. Two proteins involved in penicillin biosynthesis have PTS1 signals namely AT (ARL)⁵⁶ and phenylacetyl-CoA ligase (PCL;KI)⁵⁷. Blast analyses demonstrate that AT is only conserved in the few filamentous fungi that synthesize penicillin (*P. chrysogenum*, *A. oryzae*, *A. nidulans* and *A. flavus*). In contrast to AT, the PCL protein is conserved in all filamentous fungi. Furthermore, the *P. chrysogenum* genome encodes multiple other CoA ligases/synthetases with similarity to PCL, many of which contain a putative PTS (see **Supplementary table 7**).

PTS-signals. Of the 197 identified proteins with a putative PTS1 the most frequently observed signals are: SKL (41), AKL (41), ARL (25), SRL (16) and SKI (16), comprising 71 % of all the signals. Furthermore, a leucine residue at the carboxy terminal position appears to be preferred (in 68 % of the cases). Nevertheless, many of the identified proteins represent either hypothetical ORFs or proteins of yet unknown function. Our analysis identified only few proteins (18) with a putative PTS2, a phenomenon that is also observed in other species. This includes three separate 3-ketoacyl CoA thiolases, a typical PTS2 protein in almost all species. However, information on PTS2 sequences in filamentous fungi is scarce, implying that the ultimate PTS2 consensus may vary from that used in other species. Remarkably, in some cases genes encoding proteins with a PTS show clustering (see **Supplementary table 7**), suggesting that they have a function in the same metabolic pathway. This applies to Pc13g03340/Pc13g03350; Pc13g14410/Pc13g14420; Pc14g01040/Pc14g01070/Pc14g01090; Pc15g00410/Pc15g00420; Pc16g07060/Pc16g07070; Pc16g10060/Pc16g10070; Pc20g03610/Pc20g03630; Pc20g15650/Pc15660; Pc21g05470/Pc21g05490; Pc21g09440/Pc21g09460/Pc21g09470/Pc21g09480; Pc21g21120/Pc21g21140 and Pc22g20370/Pc22g20380. In many cases, the clustered genes show no or only weak transcription under the conditions tested. Remarkably, the Pc13g14410/Pc13g14420 genes are situated rather close to Pc13g14330, a putative cyclotetrapeptide synthetase. Additionally, the Pc22g20370/Pc22g20380 genes are located close to Pc22g20400, a putative TAF synthetase. Thus, like penicillin production, these biosynthetic routes might require microbody metabolism.

GeneChip® analysis of genomic DNA *Validation of the GeneChip®.* Fragmented DNA of *P. chrysogenum* Wisconsin54-1255 was hybridized to the DSM_PENa520255F Affymetrix GeneChip® to verify the quality of the probe set design. Nearly all (99.4%) of the probe sets designed to measure *P. chrysogenum* ORFs report a signal and are classified as ‘present’ by the GCOS software (**Supplementary Table 27**). Likewise, 99% of the probe sets designed to measure intergenic regions report a ‘present’ call, as they should when hybridizing genomic DNA. After scaling to the arithmetic mean signal of 100, most of the observed signal intensities are distributed between 50 and 200 (**Supplementary Fig. 11**) due to differences in hybridization efficiency. Most ORFs are present in only one copy on the genome. Signals that are considerably higher than 200

may suggest sequences present in the genome in multiple copies, although other techniques should be applied to establish gene copy numbers.

There are 12943 unique (so, excluding shared probe sets) annotated ORF probe sets, of which only 50 (0.4%) were called absent in this experiment. 77% of the 259 annotated ORF probe sets with a signal >200 is (partially) shared by different sequences (_s or _x suffix). Of the 651 annotated ORF probe sets that report a signal <50, 94% is a shared probe set. Most of the unique probe sets with higher signals report for tRNAs, rRNAs and some other genetic elements expected in multiple copies in the genome. 17% of these 651 probe sets is annotated as a (putative) transposon, (retro)transposable element or transposase (see **Supplementary Fig. 12** for an genome distribution overview). The 651 comprise of only a few unique probe sets for predicted ORFs, mostly hypothetical proteins, but also a putative glutamate decarboxylase GAD1 (Pc22g00970), apocytochrome b (Pc22g10800) and ribonuclease H 3-5 exonuclease like protein (Pc09g00030).

Strain comparison at genomic DNA level. In addition to the GeneChip[®] validation, a comparative analysis of the genomic DNA of the high-producing DS17690 with Wisconsin54-1255 was performed in triplicate. Although the GeneChip[®] was primarily designed for expression analysis, genomic DNA copy number changes can be detected, as long as the duplications or deletions is covered by an entire probe set, i.e. the larger part of an ORF. In practice, we restricted the analysis to detection of altered regions spanning at least several ORFs to improve the confidence in the interpretation of the measurements.

The signal intensities of the triplicate GeneChips[®] were averaged per strain and a fold change was calculated. Only regions with more than two adjacent probe sets reporting a >2-fold difference were considered in this analysis. The fold change was plotted against the genome order of the probe sets (**Supplementary fig. 13**). In total, 6 obvious and considerably large regions with different copy numbers between both strains were found, and listed in **Supplementary table 20**. Interestingly, most of the regions are flanked by one or more transposable elements, which may be indicative for the mechanisms of DNA duplication and deletion. Two gene clusters (# 1 and 6) are completely deleted from the genome of DS17690, covering in total over 50 hypothetical genes. The data indicates that the cluster with the penicillin biosynthetic genes is most likely 6- or 7-fold amplified; a more precise determination could not be made due to (technical) variation in the measurements reported by the different probe sets in the region.

Knock-out mutants The expression of all genes in DS17690 was compared to that in the Wisconsin54-1255 strain under producing conditions (+PAA). Significant changes were selected by SAM analysis with a threshold fold change larger than 2, and FDR<1%, thus identifying 1605 genes. An additional, more stringent criterium was imposed to keep only genes where the highest average signal (in either Wisconsin54-1255 or DS17690) was larger than 200. In this way, 578 genes can be selected as and these were functionally categorized according to 8 categories (**Fig. 7**).

To be able to determine the actual validity of a transcriptome-based identification of gene-products involved in β -lactam productivity, 12 genes were selected for knock-out analysis. In order to facilitate an efficient gene targeting in DS17690, a derivative with Pc22g15510, the homolog of *Saccharomyces cerevisiae* KU70, deleted⁵⁸ was used as recipient strain for the knock-out studies. Two of the 12 attempts were unsuccessful; i.e. no transformants were obtained for Pc18g01330 and Pc22g23110, respectively part of the lysine and valine pathways (**Supplementary table 21**). It is very likely that both led to auxotrophic mutants for whom growth is not supported on the minimal acetamide selection medium used. Four mutants show a significant reduction in β -lactam productivity: two genes involved in cysteine biosynthesis (Pc16g10020 and Pc22g16570), a peroxisome biogenesis factor (Pc21g04480) and the *veA* homolog (Pc13g13200). The first 3 results are in line with their (slight) increase in mRNA levels from Wisconsin54-1255 to DS17690 and under penicillinG producing conditions (+PAA). Moreover, the disturbance of peroxisome biogenesis clearly illustrates again the importance of peroxisomes in penicillin production. The *veA* mRNA is reduced with increased β -lactam productivity (**Supplementary table 21**). In *Aspergillus nidulans* VeAp was reported to repress some of the penicillin biosynthesis genes^{59,60}, so it may seem logic to find decreased mRNA levels in the DS17690 strain. Still, like in *A. nidulans*, a knock-out of the gene resulted in a severe decrease in β -lactam productivity (23.6%). As VeAp controls many more target genes, a complete silencing of the gene might very well negatively influence other factors involved in penicillin production.

One silenced gene, Pc22g13680 the putative isopenicillinN-CoA epimerase, shows an increase in β -lactam productivity, while it showed no significant change in mRNA level.. The knock-outs of the other 5 genes had no significant effect on the β -lactam productivity, suggesting that their gene products have no direct role in β -lactam biosynthesis, although their (strong) increased mRNA levels suggested otherwise. Thus,

while for some genes a clear link to β -lactam biosynthesis could be demonstrated, these results also illustrate the limitations of a solely transcriptome-based identification of targets for metabolic engineering and therefore a thorough follow-up in the form of functional analysis is needed.

SUPPLEMENTARY METHODS

Genome annotation and analysis The combined automatic and manual approach started with the analysis of all contigs larger than 5kb genes. Four different ORF predicting algorithms were applied: FGENESH⁶¹, GeneMark⁶², GENSCAN⁶³ and GeneWise⁶⁴. FGENESH, GeneMark and GENSCAN were all three run on the entire genomic sequence to provide an initial set of predicted ORFs. Preference was given to FGENESH genes. For regions without any FGENESH prediction GeneMark or GENSCAN models were extracted with preference for the GeneMark models. In addition the genomic sequence was also searched against the following three protein databases using blastx⁶⁵: a). *A. niger* proteins (Aniger)⁶⁶, b). Fungal proteins, compiled from all public available fungal sequences (Fungi), c). Non redundant protein database, compiled from all other publicly available protein sequences (PROT_NR). For all initially predicted ORFs a blastp⁶⁵ analysis against the protein databases was performed. Based on the blastp results for each gene GeneWise was run against the best blast matches. The genemodels of the initially predicted ORFs were manually adjusted in case that the blastp and GeneWise alignment indicated a suboptimal gene model.

For regions without any gene prediction with one of the three algorithms but with a significant blastx match, ORFs were manually extracted by usage of the respective GeneWise alignment. Incomplete GeneWise protein alignments were extended to the first exon upstream to the nearest start codon, and the last exon downstream to the first stop codon. For contigs smaller 5 kb only a six-frame translation was performed in order to extract all encoded exons. Transfer RNAs were identified using the tRNAScan-SE program⁶⁷. Ribosomal RNAs were identified by blastn⁶⁶ against a database of all public available rRNA sequences.

For all ORFs resulting from the above described approach an exhaustive automatic bioinformatic analysis in respect to function and structure of the respective protein was performed using the PEDANT-ProTM software⁶⁸. Annotation of description, functional categories according to the Functional Catalog (FunCat) classification

system⁶⁹, and EC numbers have been performed for each *P. chrysogenum* ORF with a multi-step semiautomatic approach:

- (1) For all known *P. chrysogenum* ORFs description, functional categories and EC numbers have been manually annotated.
- (2) *P. chrysogenum* ORFs having blastp matches against manually annotated *A. niger* ORFs⁶⁶ were assigned the respective *A. niger* annotations, if
 - a. the *A. niger* ORF was a known protein
 - b. the protein from which the *A. niger* ORF annotations were deduced from occurred also in the blast results of the *P. chrysogenum* ORF.
- (3) For *P. chrysogenum* ORFs having a blastp match (e-val $\geq 9e-04$) against either *A. niger*, Fungi, or PROT_NR database entries, annotations were assigned as follows according to the best blast match:
 - a. Best match is *A. niger* ORF with putative function: Description and FunCat transferred, e.g. "similarity to adenosine deaminase like protein An16g02890 - *Aspergillus niger*"
 - b. Best match is *A. niger* hypothetical protein: Description transferred, e.g. "strong similarity to hypothetical protein An11g00330 - *Aspergillus niger*"
 - c. Best match is fungal protein: Description transferred, e.g. "similarity to hypothetical protein 11_10_scaffold_1.tfa_290cg - *Fusarium graminearum*"
 - d. Best match against PROT_NR: Description transferred, e.g. "strong similarity to hypothetical enoyl-CoA hydratase - *Rhodopseudomonas palustris*"
- (4) All other *P. chrysogenum* ORFs having no blastp match and a protein length ≤ 79 aa, or ≥ 80 aa, were assigned as 'questionable ORF' or 'hypothetical protein', respectively.

To compare *P. chrysogenum* with other fungi, pairwise lists of orthologs were obtained using bidirectional Blastp searching of the *P. chrysogenum* predicted ORFs with ORFs of each of *Saccharomyces cerevisiae*, *Neurospora crassa*, *Magnaporthe grisea*, *Histoplasma capsulatum*, *Coccidioides immitis*, *Podospora anserina*,

Phanerochaete chrysosporium, *Gibberella zeae*, *Trichoderma reesei*, *Aspergillus nidulans*, *Aspergillus fumigatus*, *Aspergillus oryzae*, *Aspergillus terreus*, *Aspergillus flavus*, *Aspergillus clavatus* and *Aspergillus niger*. Bidirectional best hits with a blastp E-value of less than 1E-10 in both directions were considered as orthologous gene pairs.

Methods for Secondary metabolites, NJ-trees of ACVS, IPNS and siderophore

synthetases Multiple alignments to identify the putative members of gene clusters were done by ClustalW (1.81). The parameters used were: gap open penalty 10, gap extension penalty 0.05, no weight transition, hydrophilic gaps permitted, weight matrix BLOSUM.

In silico analysis of peroxisomal targeting signals *P. chrysogenum* proteins with putative microbody targeting signals (PTS1 or PTS2) were identified using the Genome 2D program⁷⁰. For identification of putative PTS1 proteins, the sequence [SCA]-[KRH]-[LMI]-COOH was used as query, consisting of the well established consensus sequence for human PTS1 proteins⁷¹ to which isoleucine was added at the C-terminal position based on the finding that *P. chrysogenum* PCL utilizes an SKI as PTS1⁷². For identification of PTS2 sequences the first 50 amino acids of each protein in the *P. chrysogenum* protein database was searched using as query: -[RK]-[LVIQ]-X-X-[LVIHQ]-[LSGAK]-X-[HQ]-[LAF]-⁷³.

Strains Independent triplicate chemostat cultivations of the *P. chrysogenum* strains Wisconsin54-1255⁷⁴ and the high producing penicillinG strain DS17690⁷⁵ were performed and monitored in the presence and absence of phenylacetic acid exactly as described before⁷⁶. DS17690 is derived from the strain improvement program of DSM after several classical mutagenesis rounds and is an offspring of the sequenced Wisconsin54-1255 strain.

Generation of knock-out mutants and β -lactam analysis The genes selected for further analysis were knocked-out in a derivative of DS17690 wherein the non-homologous end-joining pathway was deleted (by virtue of deleting the *KU70* homolog, encoded by Pc22g15510) to increase the efficiency of homologous recombination^{58,78,79}. The transcription of the selected genes was completely blocked by insertion of the *amdS* selectionmarker between the promoter and ORF of each gene. As flanking regions to

direct the targeted double homologous recombination event, 1500 basepairs of the promoter and 1500-2500 basepairs of the ORF were used (the latter depending on the length of the individual ORFs). Protoplast transformation and transformant selection was performed as described previously⁷⁹. Acetamide-consuming transformants were obtained, verified by PCR (data not shown) and sporulated. Spores were used to inoculate shake flasks with synthetic media supporting β -lactam production⁷⁹. To verify the validity of the findings two different side chain precursors were added to the media, phenyl acetic acid (PAA) and adipate (Ad), at 0.3 and 0.5 g/l respectively. Cultivation was for 168 hours at 25 °C and 280 rpm. As controls both parent strains, DS17690 and the Δ Pc22g15510 derivative, were used.

Analysis of β -lactams produced was done by quantitative ¹H NMR at 600 MHz on a Bruker Avance 600 spectrometer. To a known quantity of filtrate, a known quantity of internal standard (maleic acid), dissolved in phosphate buffer was added prior to lyophilisation. The residue was dissolved in D₂O and measured at 300 K. The delay between scans (30 s) was more than 5 times T₁ of all compounds, so the ratio between the integrals of the compounds of interest and the integral of the internal standard is an exact measure for the quantity of the β -lactams.

Media The mineral medium was set at pH 5.5 with KOH pellets and contained per liter of demineralised water 7.5 g glucose, 3.5 g (NH₄)₂SO₄, 0.8 g KH₂PO₄, 0.5 g MgSO₄•7H₂O, 10 mL of a trace element solution. The trace element solution contained 15 g•L⁻¹ Na₂EDTA•2H₂O, 0.5 g•L⁻¹ Cu₂SO₄•5H₂O, 2 g•L⁻¹ ZnSO₄•7H₂O, 2 g•L⁻¹ MnSO₄•H₂O, 4 g•L⁻¹ FeSO₄•7H₂O, and 0.5 g•L⁻¹ CaCl₂•2H₂O and was set at pH 6.0 by adding NaOH pellets. Production of penicillinG was induced by adding 0.58 g•L⁻¹ phenylacetic acid (PAA) to the medium. The appropriate amount of PAA was dissolved in demineralised water. After the pH was set to 5.5 with KOH pellets, the PAA solution was autoclaved for 40 min at 121 °C. All other components were dissolved in demineralised water and added to the PAA solution in the medium vessel via filter sterilization with a sterile 0.2 μ m capsule filter (Supor®DCF™; Pall Corporation, East Hills, USA). Glucose was heat-sterilized separately (110 °C) and added to the medium in a concentration of 7.5 g•L⁻¹. After filter sterilization and during the chemostat experiments the medium vessel was mixed continuously by a magnetic stirrer.

Chemostat cultivation Aerobic glucose-limited chemostat cultivation was performed at 25 °C in 3 - L turbine stirred bioreactors (Applikon, Schiedam, The Netherlands) with a working volume of 1.8 L. The pH was maintained at 6.5 via automated addition of 2 M NaOH (ADI 1030 biocontroller, Applikon, Schiedam, The Netherlands). The fermenter was sparged with air at a flow rate of 0.9 L•min⁻¹ using a Brooks mass-flow controller (Brooks Instruments, Hatfield, USA) and stirred at 750 rpm. The dissolved-oxygen concentration was continuously monitored with an oxygen electrode (Applisens, Schiedam, The Netherlands). Continuous cultivation was initiated after 50-60 hours of batch cultivation. The feed medium was supplied continuously by a peristaltic pump (Masterflex, Cole Parmer, USA) and the dilution rate was set at 0.03 h⁻¹ for all chemostat experiments in both the presence and absence of the penicillinG side-chain precursor phenylacetic acid (PAA). Effluent was removed discontinuously by means of a special overflow device, which has been described previously⁷⁹. The time interval between effluent removals was fixed in such a way that each time approximately 1 % of the culture volume was removed. To prevent excessive foaming, silicone antifoam (10 % vol/vol, BDH) was discontinuously added at timed intervals.

	Y_{sx}^a (g•g ⁻¹)	q_{pen}^b (μmol•g ⁻¹ •h ⁻¹)	q_{PAA} (μmol•g ⁻¹ •h ⁻¹)	q_{CO_2} (mmol•g ⁻¹ •h ⁻¹)	q_{O_2} (mmol•g ⁻¹ •h ⁻¹)
DS17690 - PAA	0.37 ± 0.01	0.00 ± 0.00	0.00 ± 0.00	1.15 ± 0.08	1.19 ± 0.11
DS17690+ PAA	0.35 ± 0.01	19.81 ± 1.47	24.04 ± 2.38	1.42 ± 0.11	1.42 ± 0.17
Wis 54-1255 - PAA	0.35 ± 0.01	0.00 ± 0.00	0.00 ± 0.00	1.45 ± 0.05	1.44 ± 0.03
Wis 54-1255 + PAA	0.35 ± 0.02	2.21 ± 0.72	35.63 ± 3.72	1.67 ± 0.13	1.73 ± 0.13

^a biomass yield on glucose (g of biomass per g of glucose consumed)

^b biomass -specific penicillinG production rate. In addition to penicillinG, intermediates and byproducts are formed, which accounts for ~6% of the consumed phenylacetic acid.

Determination of culture dry weight Culture samples (10 mL) were filtered over preweighed glass fiber filters (Type A/E, Pall Life Sciences, East Hills, USA). The filters were washed with demineralized water and dried for 20 minutes at 600 W in a microwave oven and were subsequently weighed.

Substrate and metabolite analysis Glucose concentrations in the medium were determined by HPLC analysis using an Aminex HPX-87H column (Biorad, Hercules, USA) at 60 °C with 5 mM H₂SO₄ as the mobile phase. Phenylacetic acid and penicillinG concentrations were determined by isocratic HPLC analysis using a Platinum EPS C18 column (Alltech, Deerfield, USA) at 30 °C. The mobile phase consisted of 5 M acetonitrile with 5 mM KH₂PO₄ and 6 mM H₃PO₄.

Gas analysis The exhaust gas of chemostat cultures was first passed through a condenser kept at 4 °C. The fraction of the gas that was sent to the off-gas analyzer was subsequently dried with a Perma Pure dryer (type MD-110-48P-4, Perma Pure, Toms River USA). Oxygen and carbon dioxide concentrations were determined with a NGA 2000 analyzer (Rosemount Analytical, Orville, USA). Off-gas flow rates were determined from an average of 10 measurements using a SAGA digital flow meter (Ion Science, Cambridge, UK). Specific rates of carbon dioxide and oxygen consumption were calculated as described previously⁸⁰.

GeneChip design Based on the genome sequence of Wisconsin54-1255, a proprietary GeneChip®, DSM_PENa520255F, was designed for DSM according to the Custom GeneChip® program by Affymetrix (Affymetrix, Inc., Santa Clara, CA). The chips of the 49-7875 format contain 11µm features. There are 11 pairs of 25-oligonucleotides in each probe set representing each of the approximately 13,670 annotated ORFs and supplemented with genetic elements and a selection of large intergenic sequences of *P. chrysogenum*. The probe sequences were designed based on the region in the (predicted) coding sequence stretching from 650 to 4 nt, from the 3' end. Typical Affymetrix control probe sets were included (*B. subtilis*, *E. coli*, *Bacteriophage P1* and *A. thaliana* spike controls). For several known *P. chrysogenum* genes, separate 3' end, middle and 5' end probe sets were designed that allow to monitor the efficiency of the *in vitro* transcription reaction. In addition, 1551 probe sets were included to search for the presence of yet unpredicted ORFs within large sequences (>2500nt) between two predicted ORFs or genetic elements. For this, the 'intergenic' regions were split up in 800 nt fragments (skipping 300nt from each end) and probe sets were designed for both strands (coding and reverse complement) for the sequences stretching from 300-1100 (gn3, 494 sets); 1900-2700 (gn5, 177 sets); 3500-4300 (gn7, 44 sets); 5100-5900 (gn9, 5 sets); 6700-7500 (gn11, 5 sets); 8300-9200 (gn13, 2 sets) and 10000-10800 (gn15, 2

sets) until the 300nt from the 5'end of the region. Intergenic regions on the border of two contigs were represented by two probe sets marked with A and B (e.g. gn3A, gn3B and gn5A, gn5B).

Genomic DNA isolation, labeling and hybridization Protoplasts of Wisonsin54-1255 were prepared as described⁸¹. The pellets were resuspended in 10 ml TES/SDS to lyse the cells; followed by phenol extraction and DNA precipitation⁸². RNase was degraded for 2 hours at 37 C with 0.1 mg/ml DNase-free RNase (Qiagen). Samples were again extracted with phenol and DNA was precipitated. Genomic DNA was sheared using a HydroShear® (GenomicSolutions®) in 200 µl total volume for 20 cycles. Finally, the genomic DNA was shortened to an average length of 25-100 bp using DNaseI (Sigma) and precipitated. Quality control was performed using the Bioanalyzer (Agilent).

Genomic DNA labeling and hybridization were performed at the MicroArray Division, University of Amsterdam. 15 µg purified, fragmented DNA in 36 µl 10 mM Tris pH7.5 was used for the labeling reaction. The GeneChip® DNA Labeling Reagent (Affymetrix, P/N 900542) was used to label the 3' termini of the DNA fragments, according to the Affymetrix GeneChip® Expression Analysis Technical Manual, Section 3.1.12.

Hybridisation, washing, staining and scanning were performed according to Affymetrix protocol Section 2 (eukaryotic target hybridization protocol, for 49 format).

Reproducibility was determined using two standard genes and is the average \pm standard deviation of independent cultures (n indicated in table):

	Avg CV^a	PcACTA^b	PcGDH2^c	n
DS17690 - PAA	0.21	4190 \pm 170	1240 \pm 120	3
DS17690+ PAA	0.18	3560 \pm 360	1140 \pm 270	4
Wis 54-1255 - PAA	0.10	2950 \pm 290	1030 \pm 110	3
Wis 54-1255 + PAA	0.13	3090 \pm 190	850 \pm 220	3

^a represents the average coefficient of variation (standard deviation divided by the mean) for all genes except the genes with the mean below 12

^b encoding actin; average signal and standard deviation

^c encoding glutamate dehydrogenase; average signal and standard deviation

Sampling and RNA extraction procedures 60 mL of culture broth was sampled and rapidly filtered over a glass fiber filter (Type A/E, Pall Life Sciences, East Hills, USA). The filter with mycelium was wrapped in aluminum foil, quenched in liquid nitrogen and subsequently stored at -80 °C until further use. For total RNA extraction half of the pellet was ground by mortar and pestle under constant cooling with liquid nitrogen. The powder was taken up in 5 mL of Trizol reagent (Invitrogen) and 1 mL chloroform (Sigma) and mixed well. The two phases were separated by centrifugation (4600 g, 15 minutes). Total RNA was further purified by extraction with acid-phenol/chloroform/isoamyl alcohol (5:1, pH 4.8, Ambion), followed by a chloroform extraction. Each time the phases were separated by centrifugation (4600 g, 15 minutes). Total RNA was precipitated for 30 minutes at -20°C in 96% ethanol and 0.3 M sodium acetate. After centrifugation at 23000 g for 15 minutes the RNA was dissolved in RNase free H₂O.

Probe preparation and Target hybridization Double stranded cDNA synthesis was carried out using 10 µg of total RNA and the components of the One Cycle cDNA Synthesis Kit (Affymetrix, Santa Clara, USA). The double-stranded cDNA was purified with the GeneChip® Sample Cleanup Module (Affymetrix/Qiagen) followed by in vitro transcription and labeling using the GeneChip® IVT labeling Kit (Affymetrix). Finally, labeled cRNA was purified (GeneChip® Sample Cleanup Module, Affymetrix/Qiagen) prior to fragmentation. 15 µg of fragmented, biotinylated cRNA was hybridized to the *P. chrysogenum* MicroArrays at 45 °C for 16 h as described in the Affymetrix users' manual. Washing and staining of arrays were performed using the GeneChip® Fluidics Station 400 and scanning with the Affymetrix GeneArray Scanner 3000.

Data analysis Acquisition and quantification of array images were performed using Affymetrix GeneChip® Operating Software (GCOS version 1.2). Before comparison, all arrays were globally scaled to a target value of 100 using the average signal from all gene features. To the 15,531 transcript features on the arrays, a filter was applied to extract 13,925 open reading frames. To represent the variation in triplicate measurements, the coefficient of variation (S.D. divided by the mean) was calculated. When the genes were ranked according to increasing average intensity, the average coefficient of variation showed a sharp increase for the genes with the lowest expression. Therefore, all genes in which the average expression in all conditions was below 12 were removed from the dataset. Subsequently, all remaining values below 12

were set to a value of 12. To assess differential expression, the Significance Analysis of Microarrays (SAM version 1.21) add-in to MicrosoftTM Excel was used for comparisons of replicate array experiments. The fold-change threshold and the false discovery rate values were set at 2 and 1% respectively. The data (.CEL files) were loaded into Genedata Expressionist (Genedata, Basel, Switzerland) and the genes with significantly changed expression in one of the comparisons were arranged in clusters by the *K*-means clustering tool.

Enrichment of MIPS categories was assessed for the differentially expressed genes by Fisher's Exact test employing hypergeometric distribution with a p-value cut-off of 10⁻⁴ (after Bonferroni correction). The probability was calculated as follows: the p-value of

observing *z* genes, belonging to the same functional category is:
$$P = \sum_{x=z}^{\max(N,M)} \frac{\binom{N}{x} \cdot \binom{G-N}{M-x}}{\binom{G}{M}},$$

where *N* is the total number of genes in a category, *M* is the total number of differentially expressed genes in the cluster and *G* is the total number of *P. chrysogenum* genes.

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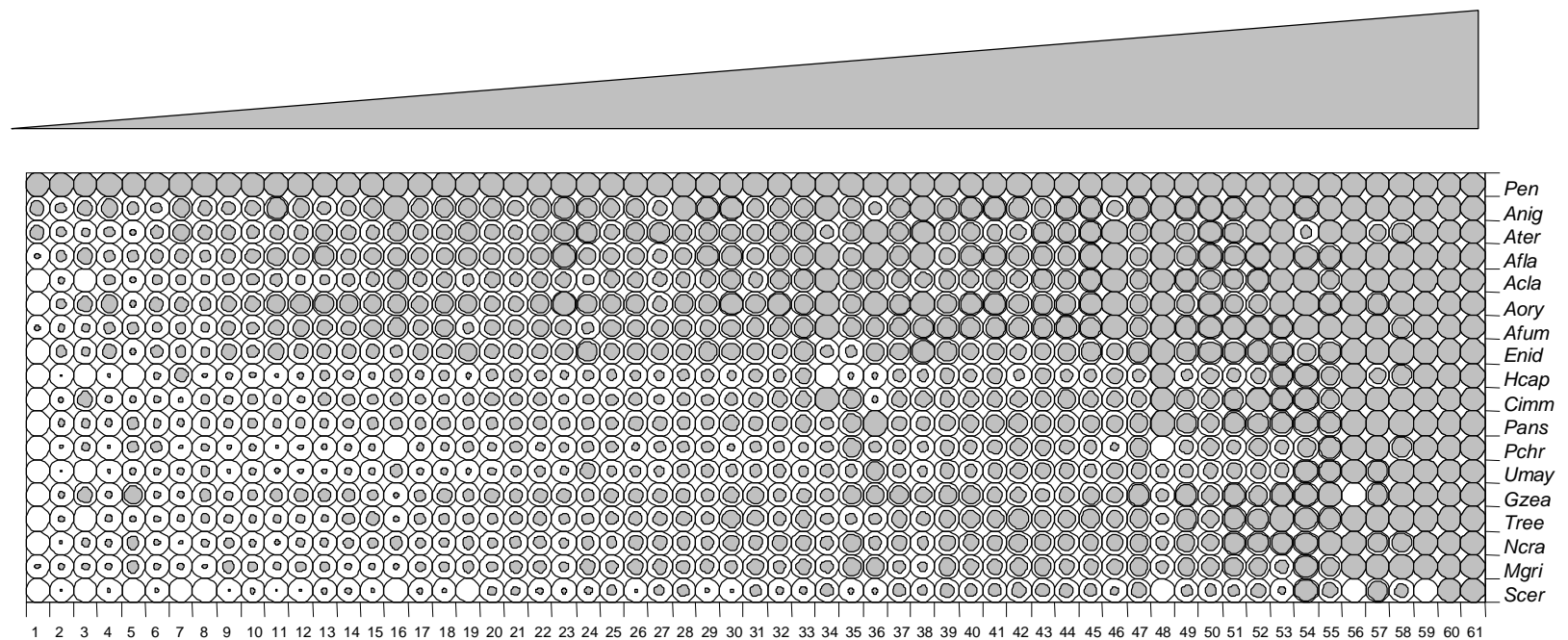
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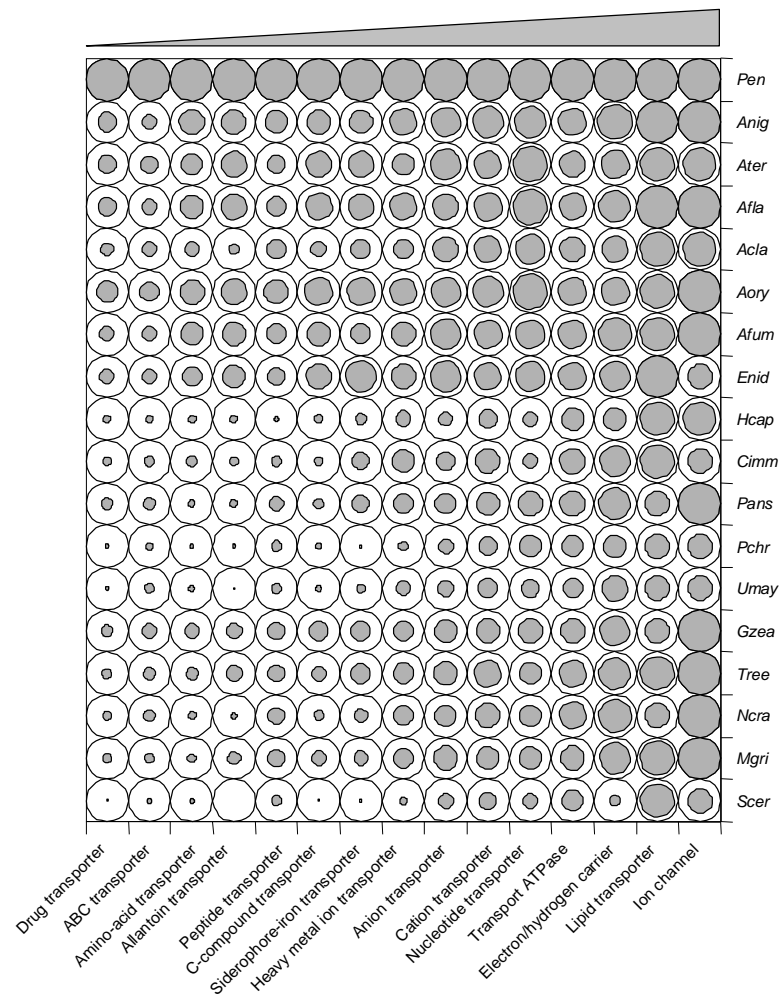
Supplementary Figure 3. Ortholog comparison of functional classes related to transporters in different fungal genomes.

Pen: *Penicillium chrysogenum*, Anig: *Aspergillus niger*, Ater: *Aspergillus terreus*, Afla: *Aspergillus flavus*, Acla: *Aspergillus clavatus*, Aory: *Aspergillus oryzae*, Afum: *Aspergillus fumigatus*, Enid: *Emericella nidulans*, Hcap: *Histoplasma capsulatum*, Cimm: *Coccidioides immitis*, Pans: *Podospira anserina*, Pchr: *Phanerochaete chrysosporium*, Umay: *Ustilago maydis*, Gzea: *Gibberella zeae*, Tree: *Trichoderma reesei*, Ncra: *Neurospora crassa*, Mgri: *Magnaporthe grisea*, Scer: *Saccharomyces cerevisiae*.

The size of the filled circles is proportional to the number of ortholog genes in each category. Categories have been sorted in respect to the number of ortholog genes over all indicated genomes increasing from left to right.

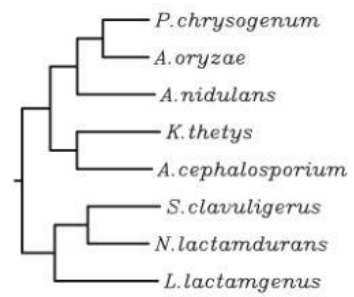


Supplementary Figure 4. Functional classification of *P. chrysogenum* specific ORFs (A), ORFs with fungal orthologs (B) and ORFs of the four non-syntenic assemblies (C). FunCat classes indicated are, 01 Metabolism, 02 Energy, 03 Cell cycle and DNA processing, 04 Transcription, 05 Protein synthesis, 06 Protein fate (folding, modification, destination), 67 Transport facilitation, 08 Cellular transport and transport mechanisms, 10 Cellular communication/Signal transduction mechanism, 11 Cell rescue, defense and virulence, 13 Regulation of/interaction with cellular environment, 40 Subcellular localization, 29 Transposable elements, 99 Unclassified proteins.

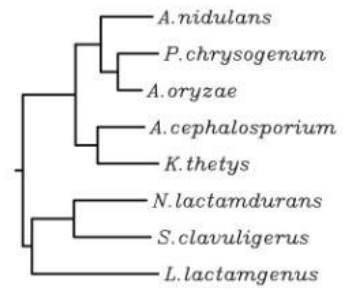


Supplementary Figure 5. NJ-tree obtained from standard ClustalW alignment of ACVS (A) and IPNS (B).

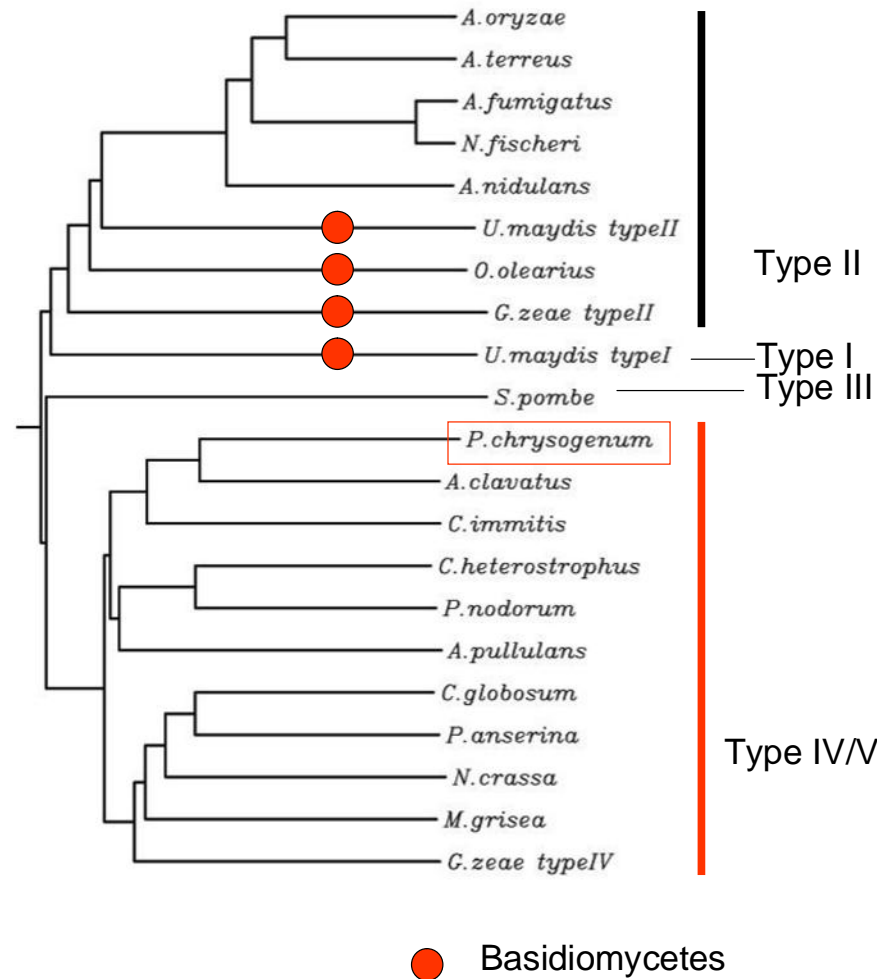
A



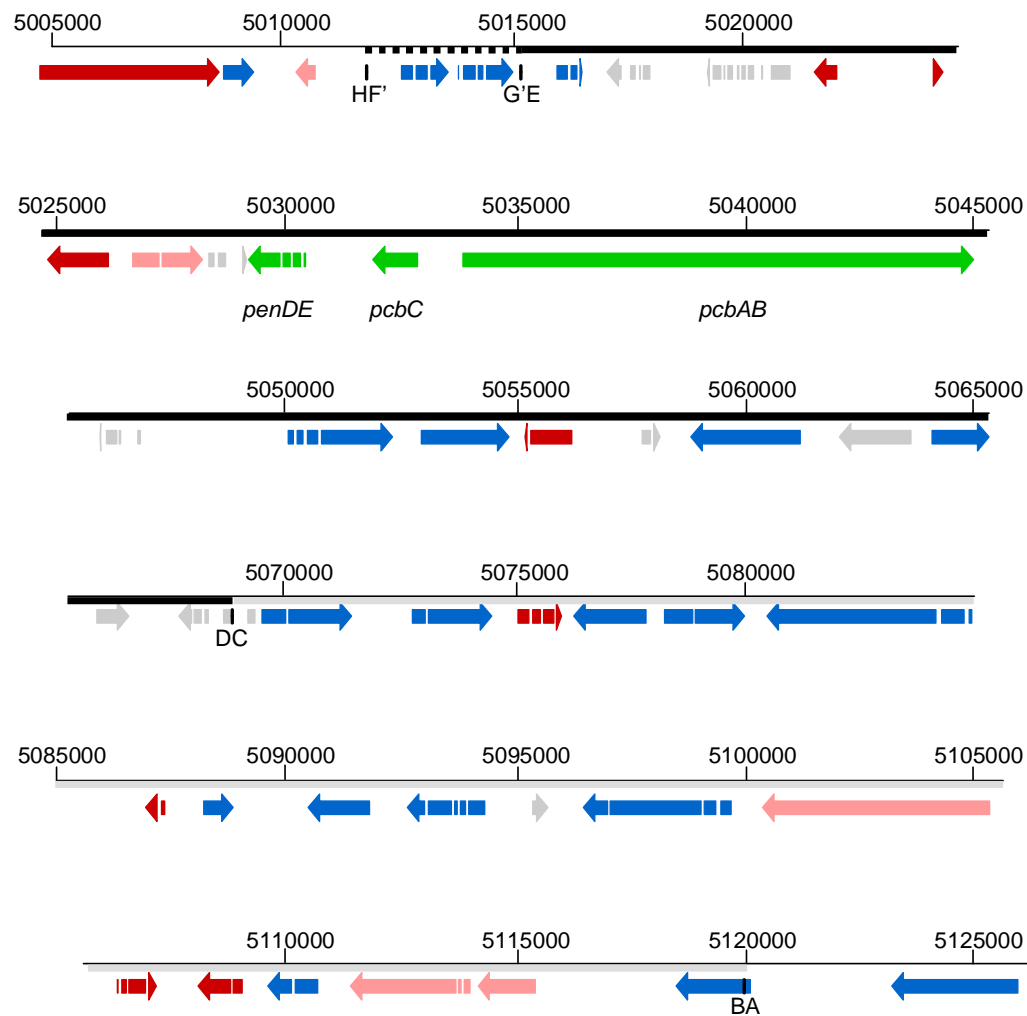
B



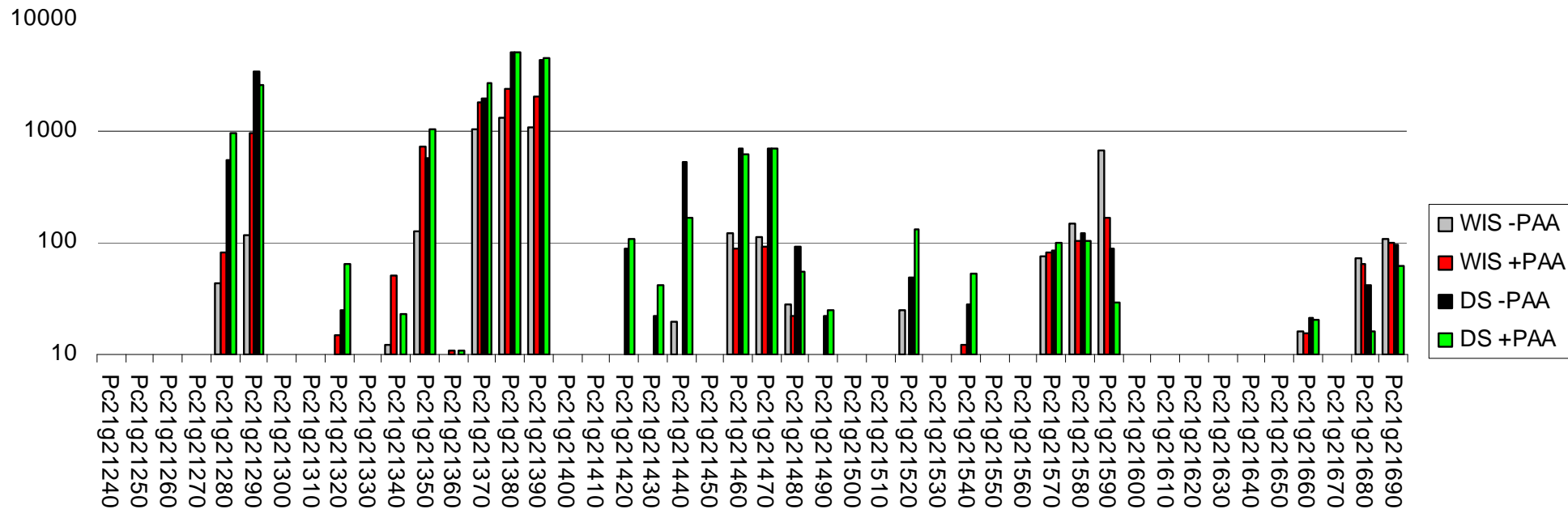
Supplementary Figure 6. NJ-tree of the known ferrichrome-type NRPS. There is an early divergence of type I/II and type III/IV/V-synthetases. Possibly two types were originally present, as still in *G. zeae* and *U. maydis*. The Ornithine N5-monooxygenase-gene is not clustered with type II although required for synthesis. Species are: *Aspergillus fumigatus*, *Aspergillus nidulans*, *Aspergillus oryzae*, *Aspergillus terreus*, *Aureobasidium pullulans*, *Chaetomium globosum*, *Cochliobolus heterostrophus*, *Gibberella zeae*, *Magnaporthe grisea*, *Neosartorya fischeri*, *Neurospora crassa*, *Omphalotus olearius*, *Podospora anserina*



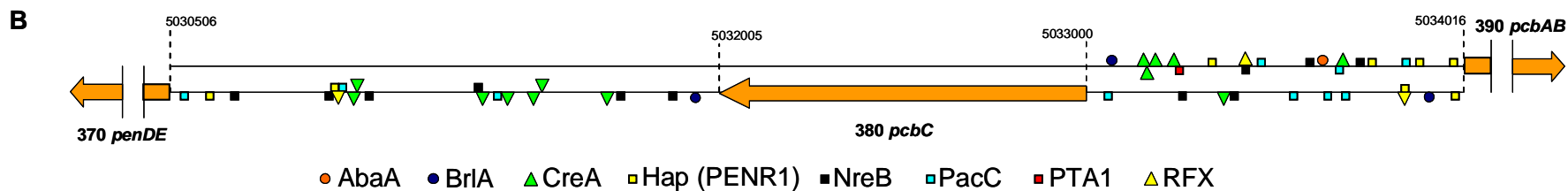
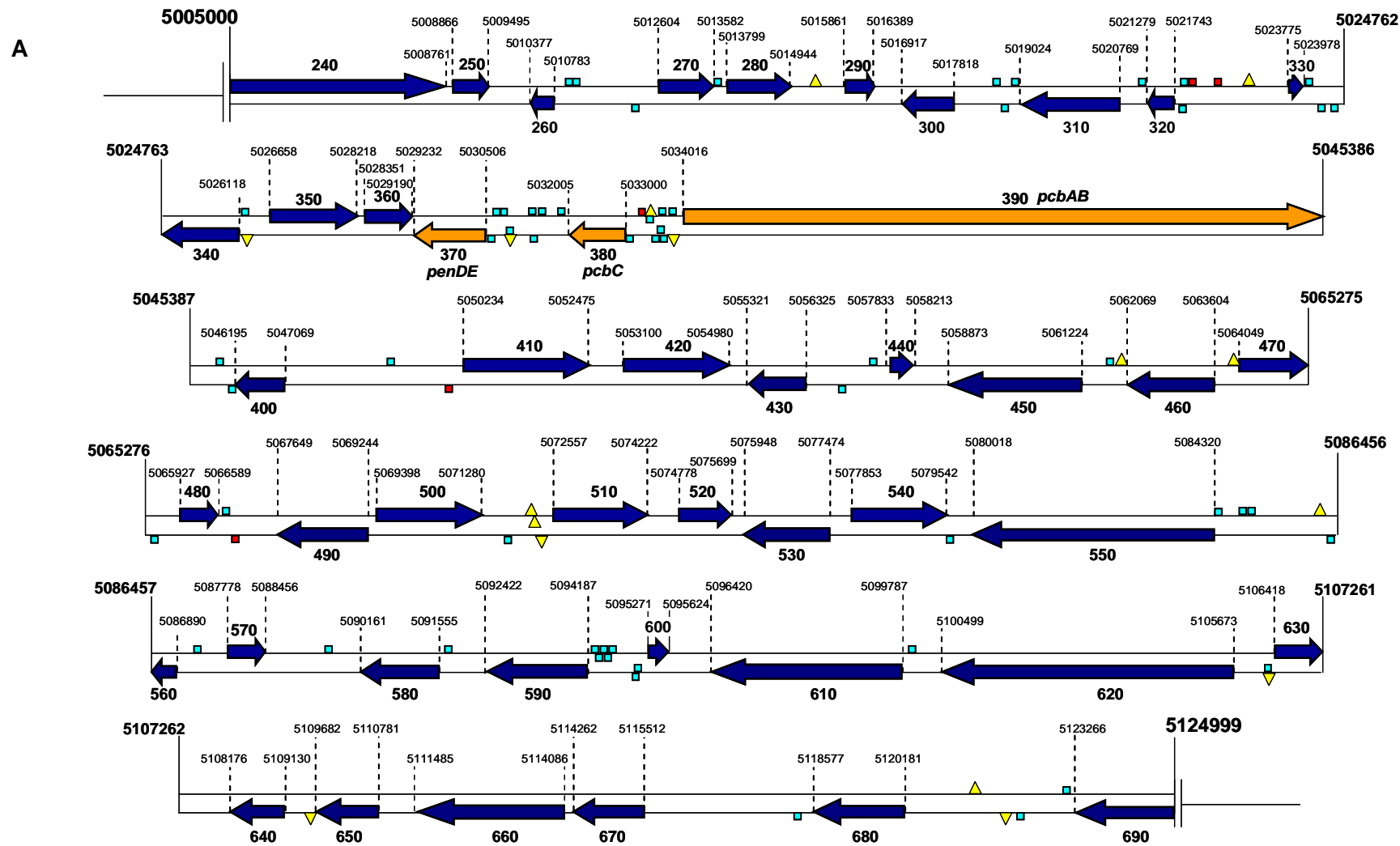
Supplementary Figure 7. Amplified chromosomal region in industrial *Penicillium* strains. Detailed overview of the region 5005000 to 5126000 from contig21 (part of chromosome I). HF', G'E, DC and AB indicate the hexamer sequences as reported¹⁷. Thick lines indicate the amplified regions: the 'flipped' region in the Wisconsin54-1255 lineage (dashed), the common part to all industrial strains (black), the part specific to the Panlabs lineage (grey). Colored arrows indicate the significance of the blastp results: hypothetical (grey), weak similarity (pink), similar (red), strong similarity (blue), known protein (green).



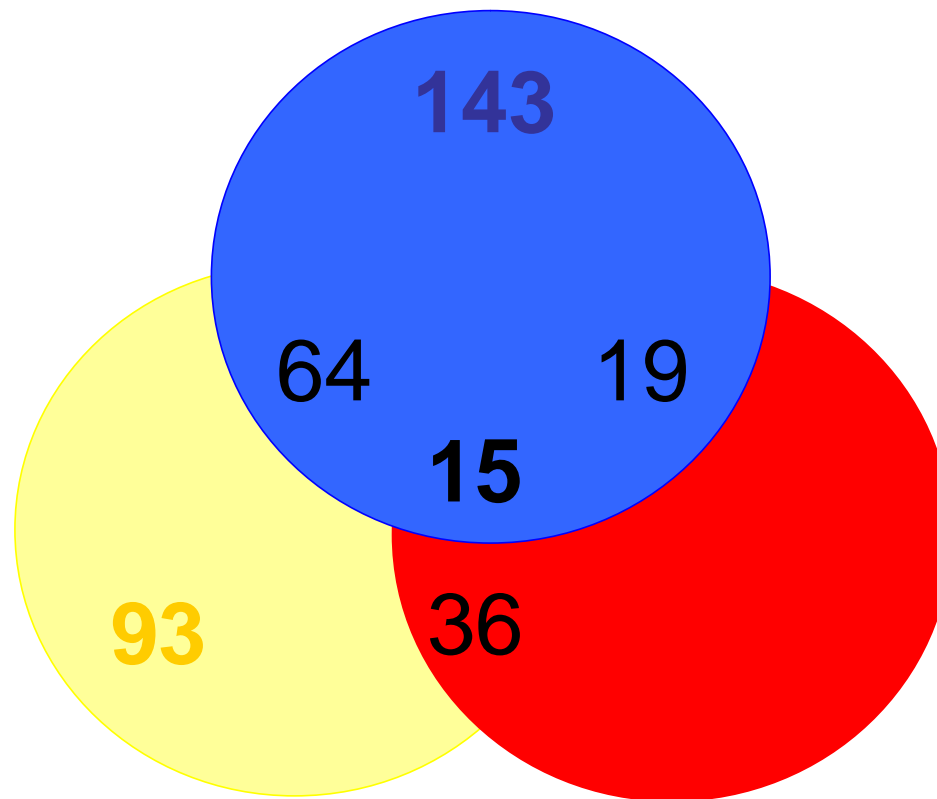
Supplementary Figure 8. Transcription profile of the amplified region in industrial *Penicillium* strains. The mean values of three individual experiments are shown (for details see **Supplementary methods**). Legend: Wisconsin54-1255 (Wis), DS17690 (DS), -PAA (no phenylacetic acid added, non-producing conditions), +PAA (phenylacetic acid added, penicillinG producing conditions).



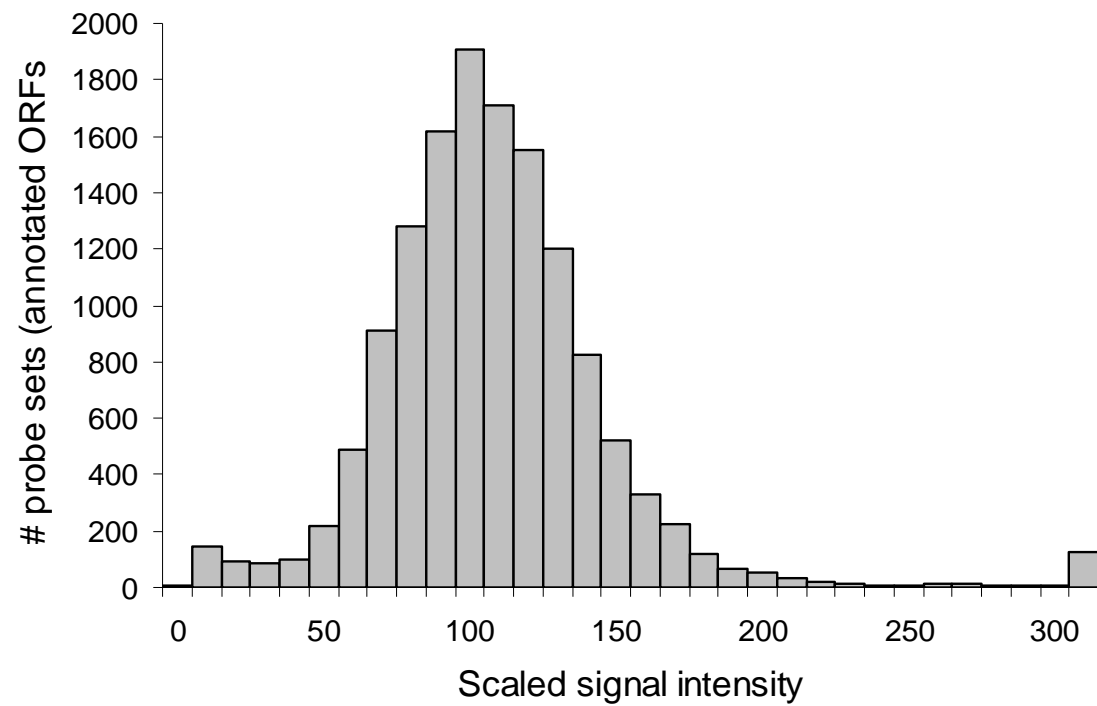
Supplementary Figure 9. DNA binding sites of different transcription factors involved in penicillin biosynthesis. A). A schematic representation of the 120-kbp sequence which contains the amplified region of *P. chrysogenum* (see **Supplementary Fig. 8**). ORFs are indicated as blue arrows, except those corresponding to the three penicillin biosynthetic genes, which are represented as orange arrows. ORFs in both strands have been numbered according to the last three numbers of the ORF code (Pc21g21xxx) and their positions on contig 21 are also indicated. The DNA binding sites present in both strands of the intergenic regions are represented for the transcription factors PacC, RFX (CpcR1-like factor) and PTA1 as blue boxes, yellow triangles and red boxes, respectively. B). Detailed schematic representation of the region including the three penicillin biosynthetic genes. Positions of the *pcbAB*, *pcbC* and *penDE* genes on contig 21 are indicated. The DNA binding regions located on the coding strand of the intergenic regions are represented as follows: AbaA (orange circle), BrlA (blue circle), CreA (green triangle), Hap (PENR1) (yellow box), NreB (black box), PacC (blue box), PTA1 (red box) and RFX (CpcR1-like factor) (yellow triangle).



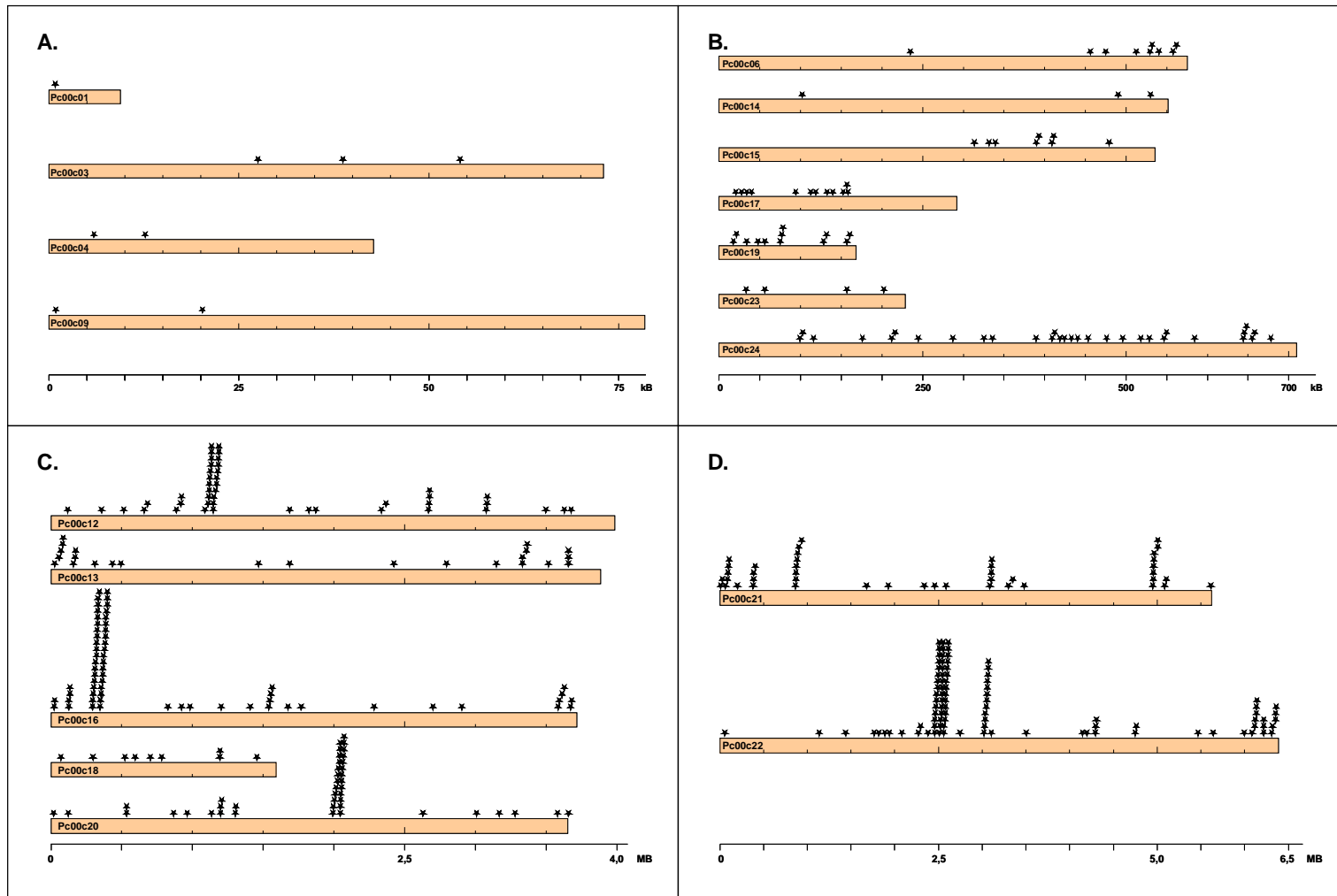
Supplementary Figure 10. Up-regulated transporter genes. Blue circle represents number of transporter genes up-regulated in the DS17690 strain with PAA versus Wisconsin54-1255 strain with PAA. Red circle represents number of genes up-regulated in the Wisconsin54-1255 strain with PAA versus Wisconsin54-1255 strain without PAA. Yellow circle represents number of genes up-regulated in the DS17690 strain with PAA versus DS17690 strain without PAA.



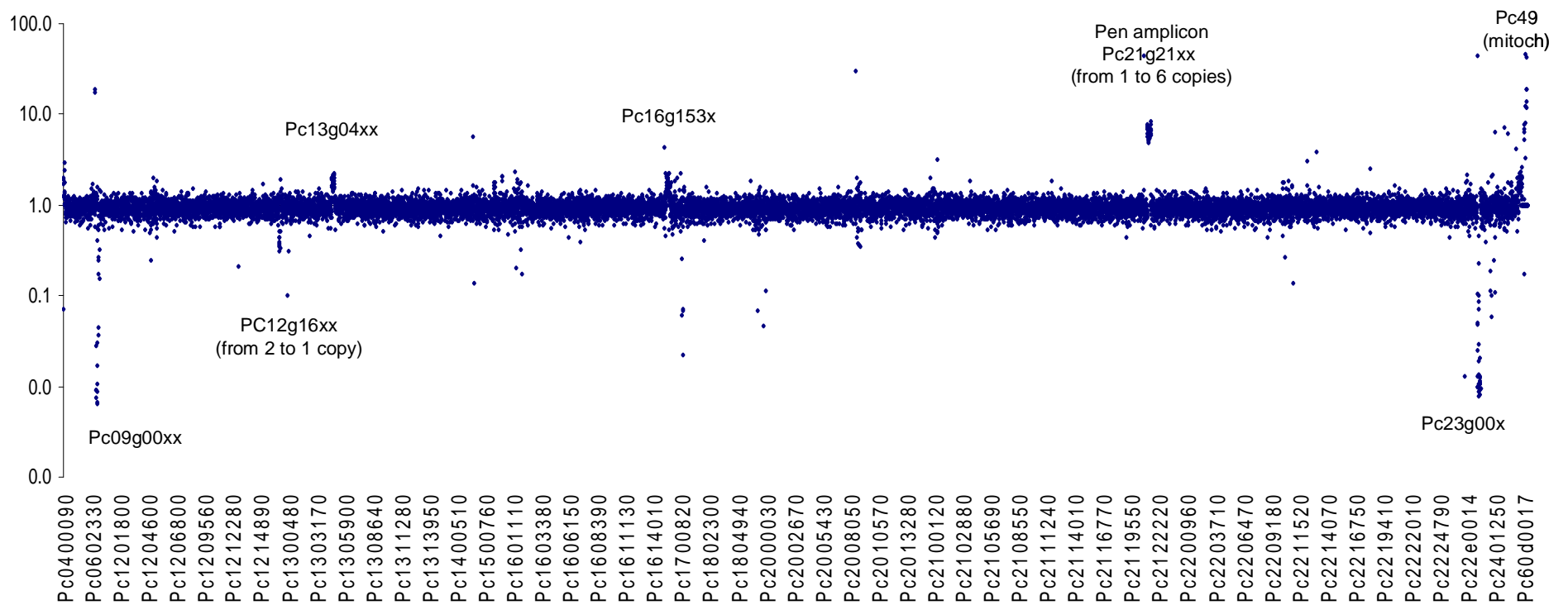
Supplementary Figure 11. Distribution of genomic DNA hybridization signal for annotated ORFs probe sets.



Supplementary Figure 12. Genome-wide comparison of genomic DNA copy number of the predicted ORFs in the production strain DS17690 vs. the reference Wisconsin54-1255. The ratios of the average intensities ($n=3$) were plotted on logarithmic scale. The indicated regions are detailed in **Supplementary table 20**.



Supplementary Figure 13. Schematic representation of genome distribution of transposon and transposon-like elements. **A.** 5kB - 100kB contigs with transposon-related sequences; **B.** 100kB - 1MB contigs with transposon-related sequences; **C.** 1MB - 5MB contigs with transposon-related sequences; **D.** >5MB contigs with transposon-related sequences. Detailed analysis of the individual elements is reported elsewhere⁸³.



Supplementary Table 23. Anticodon usage of *P. chrysogenum* tRNA genes

145 nuclear genome encoded tRNAs were identified. The number behind the amino acid indicates the number of tRNAs identified. No tRNA genes for Trp were identified. Three additional tRNA genes have been identified for which the specificity is unknown.

	.U.	.C.	.A.	.G.	
U..	Phe -	Ser 7	Tyr 1	Cys -	..U
	Phe 5	Ser -	Tyr 5	Cys 3	..C
	Leu -	Ser 1	Stop	Stop/1 ¹	..A
	Leu -	Ser 2	Stop	Trp -	..G
C..	Leu 6	Pro -	His 1	Arg 10	..U
	Leu -	Pro -	His 6	Arg -	..C
	Leu -	Pro -	Gln 3	Arg 2	..A
	Leu 3	Pro 1	Gln 5	Arg 2	..G
A..	Ile 3	Thr 7	Asn -	Ser -	..U
	Ile -	Thr -	Asn 6	Ser 4	..C
	Ile 1	Thr 3	Lys 1	Arg -	..A
	Met 6	Thr 2	Lys 6	Arg 2	..G
G..	Val 8	Ala 4	Asp -	Gly -	..U
	Val -	Ala -	Asp 7	Gly 3	..C
	Val -	Ala 2	Glu 3	Gly 3	..A
	Val 3	Ala 1	Glu 4	Gly 1	..G

¹ Sec: selenocysteine

Supplementary Table 24. *P. chrysogenum* codon usage

Numbers (%) based on all identified *P. chrysogenum* ORFs.

	.T.	.C.	.A.	.G.	
T..	Phe 35	Ser 19	Tyr 44	Cys 43	..T
	Phe 65	Ser 21	Tyr 56	Cys 57	..C
	Leu 6	Ser 15	Stop 28	Stop 37	..A
	Leu 18	Ser 16	Stop 35	Trp 100	..G
C..	Leu 19	Pro 26	His 47	Arg 18	..T
	Leu 24	Pro 27	His 43	Arg 26	..C
	Leu 11	Pro 25	Gln 44	Arg 18	..A
	Leu 22	Pro 21	Gln 56	Arg 16	..G
A..	Ile 36	Thr 25	Asn 44	Ser 12	..T
	Ile 49	Thr 34	Asn 56	Ser 18	..C
	Ile 14	Thr 24	Lys 36	Arg 12	..A
	Met 100	Thr 18	Lys 64	Arg 10	..G
G..	Val 27	Ala 28	Asp 52	Gly 27	..T
	Val 36	Ala 32	Asp 48	Gly 33	..C
	Val 11	Ala 22	Glu 43	Gly 24	..A
	Val 27	Ala 18	Glu 57	Gly 16	..G

Supplementary Table 25. Differences in *P. chrysogenum* codon usage of this study to published data

Numbers refer to the codon frequency per thousand codons. The published data are based on 97 CDS (54,967 codons). This study is based on 13,663 CDS (6,077,249 codons).

Triplet	published	this study	Diff	Triplet	published	this study	Diff	Triplet	published	this study	Diff	Triplet	published	this study	Diff
UUU	10,7	13,1	-2,4	UCU	13,5	15,7	-2,2	UAU	11,8	12,5	-0,7	UGU	4,9	6	-1,1
UUC	29,1	24,4	4,7	UCC	22,4	17,6	4,8	UAC	21,7	16	5,7	UGC	7,6	7,9	-0,3
UUA	3	5,6	-2,6	UCA	8,7	12,6	-3,9	UAA	0,7	1,1	-0,4	UGA	0,6	1,5	-0,9
UUG	14,4	16,3	-1,9	UCG	11,7	13,4	-1,7	UAG	0,5	1,4	-0,9	UGG	14,1	15	-0,9
CUU	14,3	17,2	-2,9	CCU	14	16	-2	CAU	9,3	11,8	-2,5	CGU	11,9	11,1	0,8
CUC	25,4	21,7	3,7	CCC	18,1	16,7	1,4	CAC	15	13,1	1,9	CGC	17,2	16,5	0,7
CUA	6,3	9,9	-3,6	CCA	10,6	15,3	-4,7	CAA	12,1	17,4	-5,3	CGA	6,8	11,1	-4,3
CUG	23,7	19,6	4,1	CCG	10,7	12,8	-2,1	CAG	23,9	22,3	1,6	CGG	7,5	16,5	-9
AUU	20	18,6	1,4	ACU	15,5	15	0,5	AAU	14,2	16,1	-1,9	AGU	7,5	10,1	-2,6
AUC	29,9	25,3	4,6	ACC	24,5	19,9	4,6	AAC	26,9	20,8	6,1	AGC	15,6	14,9	0,7
AUA	3,7	7,3	-3,6	ACA	10,5	14	-3,5	AAA	11,1	16,5	-5,4	AGA	4,3	7,7	-3,4
AUG	20,5	21,9	-1,4	ACG	9,8	10,4	-0,6	AAG	35,7	29,5	6,2	AGG	3,6	6,2	-2,6
GUU	17,7	16,3	1,4	GCU	24,7	22,5	2,2	GAU	25,3	28,5	-3,2	GGU	25,2	18,3	6,9
GUC	26,8	21,5	5,3	GCC	35,6	25,9	9,7	GAC	31	26,5	4,5	GGC	26,1	22	4,1
GUA	4,5	6,5	-2	GCA	14,7	18,2	-3,5	GAA	18,7	25,7	-7	GGA	16,4	15,7	0,7
GUG	19,2	16,2	3	GCG	13,3	14,9	-1,6	GAG	38	33,8	4,2	GGG	7,3	10,8	-3,5

Supplementary Table 26. Transcription Factors identified in the *P. chrysogenum* genome. *Tfam trusted matches (Domains scoring higher than the gathering threshold)

General Transcription factors involved in transcription initiation and elongation by RNA polymerases I, II and III

Transcription Factor	orf code	Description of putative <i>P. chrysogenum</i> ORF	Best blast homolog to putative <i>P. chrysogenum</i> ORF		Average transcript levels @				Transcription factor Domain*
			Species	Gene code	e-value	WIS - PAA	WIS + PAA	DS - PAA	DS + PAA
BDF1	Pc22g07210	strong similarity to transcription factor Bdf1 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	SS6956	1e-69	373.8	342.2	261.2	215.6
	Pc22g1540	strong similarity to RNA polymerase I transcription factor RRN3 - <i>Homo sapiens</i>	<i>Homo sapiens</i>	RRN3	3e-96	219.9	219.9	157.3	157.3
	RRN6	weak similarity to transcription initiation factor Rrn6 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	RRN6	1e-11	51.4	40.7	29.9	17.5
SPBC336.07	Pc12g101450	similarity to hypothetical transcription factor subunit SPBC336.07 - <i>Schizosaccharomyces pombe</i>	<i>Schizosaccharomyces pombe</i>	SPBC336.07	7e-20	84.7	54.1	70.5	60.5
	Pc16g13800	strong similarity to large subunit of transcription factor TFIIA like protein Arp112770 - <i>Aspergillus niger</i>	<i>Aspergillus niger</i>	Arp112770	4e-25	74.8	48.5	41.7	31.1
	Pc21g02570	strong similarity to small subunit of transcription factor TFIIA Tox2 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	Tox2	1e-13	34.2	36.1	25.8	33.1
TFIIB	Pc18g151520	strong similarity to transcription initiation factor TFIIB subunit - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	TFIIB	1e-90	813.3	386.2	728.7	679.8
	Pc18g10320	strong similarity to TFIIB related factor hBRF - <i>Homo sapiens</i>	<i>Homo sapiens</i>	hBRF	5e-84	26.6	29.8	20.2	14.0
	Pc21g12070	strong similarity to transcription initiation factor IIB chain like protein Arf1500960 - <i>Aspergillus niger</i>	<i>Aspergillus niger</i>	Arf1500960	1e-20	83.9	69.1	76.9	61.3
TFIIC	Pc13g10100	strong similarity to RNA polymerase II transcription initiation factor TFIIC subunit like protein Arf04310 - <i>Aspergillus niger</i>	<i>Aspergillus niger</i>	Arf04310	1e-20	813.3	386.2	728.7	679.8
	Pc13g10840	strong similarity to hypothetical transcription factor IIC-like protein CAE47090.1 - <i>Aspergillus fumigatus</i>	<i>Aspergillus fumigatus</i>	CAE47090.1	1e-11	69.5	52.5	40.4	23.1
	Pc16g09970	strong similarity to assembling subunit of transcription factor IIC - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	IIC	1e-15	128.1	111.3	104.2	104.2
TFIID (TBP and TBP Associated Factors; TAFs)	Pc12g15530	strong similarity to transcription factor IIC 95K chain Tlc1 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	Tlc1	1e-25	26.6	22.0	13.3	12.3
	Pc12g01100	similarity to component of the TBP-associated protein complex TAF47 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	TAF47	1e-11	168.0	175.1	137.6	107.5
	Pc12g11110	strong similarity to subunit of transcription initiation factor TFIID pTf1p - <i>Schizosaccharomyces pombe</i>	<i>Schizosaccharomyces pombe</i>	pTf1p	1e-20	84.7	54.1	70.5	60.5
TFIIE	Pc12g11400	strong similarity to subunit of transcription initiation factor TFIIE like protein Arf150530 - <i>Aspergillus niger</i>	<i>Aspergillus niger</i>	Arf150530	1e-20	813.3	386.2	728.7	679.8
	Pc13g10670	strong similarity to subunit of transcription initiation factor TFIIE TATA-box binding protein TBP - <i>Aspergillus nidulans</i>	<i>Aspergillus nidulans</i>	TBP	1e-20	813.3	386.2	728.7	679.8
	Pc18g02510	strong similarity to subunit of transcription initiation factor TFIIE TFIIE subunit TAF19 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	TAF19	1e-11	69.5	52.5	40.4	23.1
TFIIF	Pc12g14080	strong similarity to subunit of transcription initiation factor TFIIF TFIIF subunit TAF19 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	TAF19	1e-11	69.5	52.5	40.4	23.1
	Pc21g10490	strong similarity to subunit of transcription initiation factor TFIIF Tsm1 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	Tsm1	1e-11	69.5	52.5	40.4	23.1
	Pc21g10240	strong similarity to subunit of transcription initiation factor TFIIF TFIIF subunit TAF19 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	TAF19	1e-11	69.5	52.5	40.4	23.1
TFIIG	Pc12g16340	strong similarity to subunit of transcription initiation factor TFIIG TFIIG subunit TAF19 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	TAF19	1e-11	69.5	52.5	40.4	23.1
	Pc21g12800	strong similarity to subunit of transcription initiation factor TFIIG TFIIG subunit TAF19 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	TAF19	1e-11	69.5	52.5	40.4	23.1
	Pc21g12920	strong similarity to subunit of transcription initiation factor TFIIG TFIIG subunit TAF19 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	TAF19	1e-11	69.5	52.5	40.4	23.1
TFIIR	Pc22g25300	strong similarity to subunit of transcription initiation factor TFIIR TFIIR subunit TAF19 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	TAF19	1e-11	69.5	52.5	40.4	23.1
	Pc12g02300	strong similarity to alpha subunit of transcription initiation factor TFIIR TFIIR subunit TAF19 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	TAF19	1e-11	69.5	52.5	40.4	23.1
	Pc12g03030	strong similarity to beta subunit of transcription initiation factor TFIIR TFIIR subunit TAF19 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	TAF19	1e-11	69.5	52.5	40.4	23.1
TFIIS	Pc12g07600	strong similarity to alpha subunit of transcription initiation factor TFIIS TFIIS subunit TAF19 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	TAF19	1e-11	69.5	52.5	40.4	23.1
	Pc12g07350	strong similarity to small subunit of transcription initiation factor TFIIS TFIIS subunit TAF19 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	TAF19	1e-11	69.5	52.5	40.4	23.1
	Pc12g16480	weak similarity to subunit of transcription initiation factor TFIIS TFIIS subunit TAF19 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	TAF19	1e-11	69.5	52.5	40.4	23.1
TFIIR	Pc13g06540	strong similarity to subunit of transcription initiation factor TFIIR TFIIR subunit TAF19 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	TAF19	1e-11	69.5	52.5	40.4	23.1
	Pc12g07740	strong similarity to subunit of transcription initiation factor TFIIR TFIIR subunit TAF19 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	TAF19	1e-11	69.5	52.5	40.4	23.1
	Pc16g12370	strong similarity to subunit of transcription initiation factor TFIIR TFIIR subunit TAF19 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	TAF19	1e-11	69.5	52.5	40.4	23.1
TFIIR	Pc21g06450	strong similarity to subunit of transcription initiation factor TFIIR TFIIR subunit TAF19 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	TAF19	1e-11	69.5	52.5	40.4	23.1
	Pc12g04800	strong similarity to subunit of transcription initiation factor TFIIR TFIIR subunit TAF19 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	TAF19	1e-11	69.5	52.5	40.4	23.1
	Pc21g11600	strong similarity to subunit of transcription initiation factor TFIIR TFIIR subunit TAF19 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	TAF19	1e-11	69.5	52.5	40.4	23.1
Transcription elongation factor	Pc12g08440	weak similarity to subunit of transcription initiation factor TFIIR TFIIR subunit TAF19 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	TAF19	1e-11	69.5	52.5	40.4	23.1
	Pc13g04600	strong similarity to transcription elongation factor like protein Arf0500780 - <i>Aspergillus niger</i>	<i>Aspergillus niger</i>	Arf0500780	1e-20	813.3	386.2	728.7	679.8
	Pc15g15070	strong similarity to transcription elongation factor TFIIS encoded by tfs1p - <i>Schizosaccharomyces pombe</i>	<i>Schizosaccharomyces pombe</i>	tfs1p	1e-20	84.7	54.1	70.5	60.5
Transcription elongation factor	Pc12g03610	strong similarity to transcription elongation protein SpT5 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	SpT5	1e-11	69.5	52.5	40.4	23.1
	Pc21g08080	strong similarity to transcription elongation protein SpT6 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	SpT6	1e-11	69.5	52.5	40.4	23.1
	Pc21g11690	weak similarity to transcription elongation protein SpT5 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	SpT5	1e-11	69.5	52.5	40.4	23.1
Transcription elongation factor	Pc12g17890	similarity to protein SpT4 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	SpT4	1e-11	69.5	52.5	40.4	23.1

Specific Transcription Factors

Transcription Factor	orf code	Description of putative <i>P. chrysogenum</i> ORF	Best blast homolog to putative <i>P. chrysogenum</i> ORF		Average transcript levels @				Transcription factor Domain*		
			Species	Gene code	e-value	WIS -PAA	WIS +PAA	DS -PAA		DS +PAA	
ABCA1	Pc16g09610	strong similarity to protein AbaA - <i>Aspergillus nidulans</i>	<i>Aspergillus nidulans</i>	AbA	0.0	22.6	8.3	63.5	4.3	TEA/ATTS	
	Pc20g06400	strong similarity to Cys2-His2 zinc finger transcription factor ACE1 - <i>Hypocrea jecorina</i>	<i>Neurospora crassa</i>	ACE1	1e-165	88.7	77.8	61.2	37.3	C2H2	
	Pc20g02020	weak similarity to transcription factor involved in acriflavine resistance acr-2 - <i>Neurospora crassa</i> [putative sequencing error]	<i>Neurospora crassa</i>	BX146200, 46	2e-46	68.4	57.4	63.9	39.0	Zn(II)-Cys6	
	Pc20g01550	strong similarity to transcription factor involved in acriflavine resistance acr-2 - <i>Neurospora crassa</i>	<i>Neurospora crassa</i>	gene: "B11E5.460"; product: "putative protein";	BX426200, 46	5e-10	64.8	67.4	51.1	Zn(II)-Cys6	
	Pc12g08760	weak similarity to transcription factor involved in acriflavine resistance acr-2 - <i>Neurospora crassa</i>	<i>Neurospora crassa</i>	gene: "17E5.150"; product: "conserved hypothetical protein";	NC17E5, 14	3e-18	5.4	7.3	7.5	1.4	No domain identified
ACR-2	Pc13g08450	weak similarity to transcription factor involved in acriflavine resistance acr-2 - <i>Neurospora crassa</i>	<i>Neurospora crassa</i>	gene: "B11E5.160"; product: "putative protein";	BX426200, 46	1e-12	86.4	88.9	89.9	89.9	Zn(II)-Cys6
	Pc16g12430	strong similarity to transcription factor involved in acriflavine resistance acr-2 - <i>Neurospora crassa</i>	<i>Neurospora crassa</i>	acr-2 protein - <i>Neurospora crassa</i>	ST5237	3e-18	67.1	79.8	63.0	67.6	Zn(II)-Cys6
	Pc17g00670	weak similarity to transcription factor involved in acriflavine resistance acr-2 - <i>Neurospora crassa</i>	<i>Neurospora crassa</i>	gene: "acr-2"; <i>Neurospora crassa</i> acr-2 DNA, complete cds.	NC6893, 3	3e-24	104.9	109.0	113.5	96.5	Zn(II)-Cys6
	Pc18g00590	strong similarity to transcription factor involved in acriflavine resistance acr-2 - <i>Neurospora crassa</i>	<i>Neurospora crassa</i>	acr-2 protein - <i>Neurospora crassa</i>	ST5237, 3	9e-12	50.4	46.6	46.6	51.7	Zn(II)-Cys6
	Pc20g11000	weak similarity to transcription factor involved in acriflavine resistance acr-2 - <i>Neurospora crassa</i> [putative sequencing error]	<i>Neurospora crassa</i>	gene: "B7F21.070"; product: "hypothetical protein";	NC78F21, 6	3e-04	9.1	7.1	13.0	16.5	No domain identified
	Pc20g11950	weak similarity to transcription factor involved in acriflavine resistance acr-2 - <i>Neurospora crassa</i>	<i>Neurospora crassa</i>	zinc finger protein 1 homolog [imported] - <i>Neurospora crassa</i>	149729	9e-09	81.0	80.7	53.4	54.9	Zn(II)-Cys6
	Pc21g04570	weak similarity to transcription factor involved in acriflavine resistance acr-2 - <i>Neurospora crassa</i>	<i>Neurospora crassa</i>	gene: "B11E5.460"; product: "putative protein";	BX426200, 46	2e-21	244.6	260.7	203.5	193.2	Zn(II)-Cys6
	Pc21g09380	weak similarity to transcription factor involved in acriflavine resistance acr-2 - <i>Neurospora crassa</i>	<i>Neurospora crassa</i>	gene: "B7F21.070"; product: "hypothetical protein";	NC78F21, 6	1e-12	15.6	16.1	22.5	29.4	Zn(II)-Cys6
	Pc22g21330	weak similarity to transcription factor involved in acriflavine resistance acr-2 - <i>Neurospora crassa</i>	<i>Neurospora crassa</i>	OB-schizothecio DNA topoisomerase II-related; - <i>Escherichia coli</i>	NC12E1, 1	1e-11	123.5	121.5	20.2	19.6	ADN Zn finger, Bacterial regulatory helix-turn-helix proteins (AraC family)
	Pc12g17410	strong similarity to ADA and SAGA histone acetyltransferase subunit Ada3 - <i>Escherichia coli</i>	<i>Escherichia coli</i>	gene: "ada3"; product: "ada regulatory protein";	AD014348, 9	2e-27	27.7	28.1	29.3	44.5	Ada Zn finger, Bacterial regulatory helix-turn-helix proteins (AraC family)
ADA2, ADA3	Pc16g09180	strong similarity to ADA and SAGA histone acetyltransferase subunit Ada3 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	gene: "ADA3"; product: "alteration/deficiency in activation 3 protein";	SCAD43A, 1	8e-32	31.6	18.0	9.7	10.1	No domain identified
	Pc22g05790	strong similarity to transcription adaptor Ada2 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	gene: "GSD.16.020"; product: "related to transcription adaptor ADA2";	SPR13808, 2	1e-167	254.5	259.0	119.6	104.7	ZZ zinc finger Myb DNA binding domain, SWIRM
ADR1	Pc22g05790	strong similarity to transcription activator ADR1 - <i>Saccharomyces cerevisiae</i>	<i>Saccharomyces cerevisiae</i>	gene: "SPR13808.2"; product: "transcription activator ADR1";	SPR13808, 2	1e-167	254.5	259.0	119.6	104.7	Abn N-terminal Hsp90 ATPase, NARMH, AHSA1 (Activator of Hsp90)
	Pc13g10310	strong similarity to sterigmatocystin synthesis transcription regulator ADR1 - <i>Aspergillus nidulans</i>	<i>Aspergillus nidulans</i>	gene: "GSP.121.100"; product: "sterigmatocystin synthetase";	AF549411, 1	1e-06	64.5	63.9	66.0	91.2	Zn(II)-Cys6
AFLR	Pc16g02310	weak similarity to aflatoxin biosynthesis regulatory protein aflR - <i>Aspergillus parasiticus</i>	<i>Aspergillus parasiticus</i>	transcription regulator, nuclear cluster zinc finger protein - fission	T39467, 4	4e-05	144.2	130.1	125.5	94.4	Zn(II)-Cys6
	Pc16g02830	strong similarity to aflatoxin biosynthesis regulatory protein aflR - <i>Aspergillus parasiticus</i>	<i>Aspergillus parasiticus</i>	gene: "aflR"; product: "AFLR: Aspergillus flavus isolate BNA04 AFLR	AF1460, 1	2e-04	10.5	6.4	11.1	9.8	Zn(II)-Cys6
	Pc16g02740	strong similarity to aflatoxin biosynthesis regulatory protein aflR - <i>Aspergillus flavus</i>	<i>Aspergillus flavus</i>	gene: "aflR"; product: "AFLR: Aspergillus nomius isolate AC50 AFLR	AF441406, 1	8e-09	145.2	162.3	200.6	196.3	Zn(II)-Cys6
	Pc20g15060	strong similarity to aflatoxin biosynthesis regulatory protein aflR - <i>Aspergillus flavus</i>	<i>Aspergillus flavus</i>	gene: "aflR"; product: "AFLR: Aspergillus nomius isolate BNA103P	AF441421, 1	6e-05	32.3	26.4	47.1	70.4	C2H2, Zn(II)-Cys6
	Pc12g04750	strong similarity to aflatoxin biosynthesis regulatory protein aflR - <i>Aspergillus parasiticus</i>	<i>Aspergillus parasiticus</i>	gene: "AKA12H2.19c"; product: "hypothetical protein";	AF412H2, 19	9e-68	68.6	44.0	50.7	39.3	Zn(II)-Cys6, Fungal specific transcription factor domain
ALCR	Pc22g07530	strong similarity to sterigmatocystin synthesis transcription regulator aflR - <i>Aspergillus nidulans</i>	<i>Aspergillus nidulans</i>	transcription regulator, nuclear cluster zinc finger protein - fission	AF412H2, 19	9e-68	68.6	44.0	50.7	39.3	Zn(II)-Cys6
	Pc12g2800	strong similarity to transcription regulator ALCR - <i>Aspergillus nidulans</i>	<i>Aspergillus nidulans</i>	REGULATORY PROTEIN ALCR - <i>Aspergillus nidulans</i>	ALCR, EMENI	0.0	108.0	123.8	142.8	236.0	Zn(II)-Cys6
	Pc06g01210	strong similarity to zinc-finger transcription factor amda - <i>Aspergillus nidulans</i>	<i>Aspergillus nidulans</i>	gene: "MG1"; product: "MG repressor"; Candida utilis MG1 gene for	CU277830, 1	4e-07	1.9	2.6	2.1	1.0	C2H2
	Pc12g12000	strong similarity to zinc-finger transcription factor amda - <i>Aspergillus nidulans</i>	<i>Aspergillus nidulans</i>	DNA-binding protein amda - <i>Emicella nidulans</i>	SE1908	6e-11	35.5	74.4	40.1	53.7	C2H2
	Pc16g09130	strong similarity to zinc-finger transcription factor amda - <i>Aspergillus nidulans</i>	<i>Aspergillus nidulans</i>	similarity: "Ste12-like transcription factor" - <i>Neurospora crassa</i> Ste12-like	AF1460, 1	2e-04	10.5	6.4	11.1	9.8	C2H2
AMDA	Pc16g16880	strong similarity to zinc-finger transcription factor amda - <i>Aspergillus nidulans</i>	<i>Aspergillus nidulans</i>	DNA-binding protein amda - <i>Emicella nidulans</i>	SE1908	5e-95	43.5	2.5	6.2	12.2	C2H2
	Pc20g06540	strong similarity to zinc-finger transcription factor amda - <i>Aspergillus nidulans</i>	<i>Aspergillus nidulans</i>	Aspergillus fumigatus regA gene for transcription factor RegA,	AJ82911, 1	8e-64	42.1	45.2	64.6	121.8	Zn(II)-Cys6
	Pc12g10180	weak similarity to zinc-finger transcription factor amda - <i>Aspergillus nidulans</i>	<i>Aspergillus nidulans</i>	DNA-binding protein amda - <i>Emicella nidulans</i>	SE1908	4e-28	539.9	420.1	258.9	249.2	C2H2
	Pc12g17100	strong similarity to zinc-finger transcription factor amda - <i>Aspergillus nidulans</i>	<i>Aspergillus nidulans</i>	strong binding protein amda - <i>Emicella nidulans</i>	SE1908	1e-60	46.0	19.7	46.2	69.5	C2H2
	Pc12g17480	strong similarity to zinc-finger transcription factor amda - <i>Aspergillus nidulans</i>	<i>Aspergillus nidulans</i>	DNA-binding protein amda - <i>Emicella nidulans</i>	SE1908	0.0	45.6	40.9	59.1	83.9	C2H2
AMDR	Pc12g21660	strong similarity to zinc-finger transcription factor amda - <i>Aspergillus nidulans</i>	<i>Aspergillus nidulans</i>	DNA-binding protein amda - <i>Emicella nidulans</i>	SE1908	7e-89	16.4	15.8	21.2	20.3	C2H2
	Pc12g24840	strong similarity to zinc-finger transcription factor amda - <i>Aspergillus nidulans</i>	<i>Aspergillus nidulans</i>	strong binding protein amda - <i>Emicella nidulans</i>	SE1908	2e-14	33.5	48.9	54.0	64.0	C2H2
	Pc22g02870	strong similarity to zinc-finger transcription factor amda - <i>Aspergillus nidulans</i>	<i>Aspergillus nidulans</i>	DNA-binding protein amda - <i>Emicella nidulans</i>	SE1908	0.0	214.9	235.5	214.1	206.1	Zn(II)-Cys6, Fungal specific transcription factor domain
	Pc16g06250	strong similarity to transcription regulator amdr - <i>Aspergillus nidulans</i>	<i>Aspergillus nidulans</i>	gene: "amdr"; product: "regulatory protein"; Aspergillus oryzae	AOAMR, 1	0.0	214.9	235.5	214.1	206.1	Zn(II)-Cys6, Fungal specific transcription factor domain
	Pc12g15680	strong similarity to DNA binding regulatory protein amdr - <i>Aspergillus nidulans</i>	<i>Aspergillus nidulans</i>	gene: "RGAM"; product: "proline-rich zinc finger protein"; <i>S.cerevisiae</i>	SCRGAM, 1	1e-11	137.0	104.1	83.7	40.0	C2H2
AMDX	Pc20g04780	strong similarity to DNA binding regulatory protein amdr - <i>Aspergillus nidulans</i>	<i>Aspergillus nidulans</i>	DNA binding regulatory protein - <i>Emicella nidulans</i>	EMENI, 1	1e-03	174.0	174.0	174.0	174.0	C2H2
	Pc12g19890	strong similarity to DNA binding regulatory protein amdx - <i>Aspergillus nidulans</i>	<i>Aspergillus nidulans</i>	DNA binding regulatory protein - <i>Emicella nidulans</i>	TR1803, 0	0.0	195.7	269.1	179.8	215.6	C2H2
	Pc12g04420	strong similarity to transcription regulator of maltose utilization amyR - <i>Aspergillus oryzae</i>	<i>Aspergillus oryzae</i>	gene: "10AC10.220"; product: "related to transcription activator amyR";	NC10410, 21	6e-23	176.1	156.2	137.9	156.8	Zn(II)-Cys6, Fungal specific transcription factor domain
	Pc12g08700	strong similarity to transcription regulator of maltose utilization amyR - <i>Aspergillus oryzae</i>	<i>Aspergillus oryzae</i>	gene: "facB"; product: "FacB; Aspergillus niger putative DNA binding	ANL55039, 1	3e-07	69.5	65.7	138.1	148.2	Zn(II)-Cys6, Fungal specific transcription factor domain
	Pc12g19890	strong similarity to transcription regulator of maltose utilization amyR - <i>Aspergillus oryzae</i>	<i>Aspergillus oryzae</i>	gene: "facB"; product: "FacB; Aspergillus niger putative DNA binding	ANL55039, 1	3e-07	69.5	65.7	138.1	148.2	Zn(II)-Cys6, Fungal specific transcription factor domain

AMVR	Pc13g12960	weak similarity to transcription regulator of maltose utilization amyR - Aspergillus oryzae	gene: "CAT8", product: "Cat8", Candida albicans Ngt1 (NTG1) gene.	AF222908.3	4e-08	11.0	11.0	25.4	12.9	Zn(II)Cys6
	Pc16g06580	weak similarity to transcription regulator of maltose utilization amyR - Aspergillus oryzae	gene: "H8F11.070", product: "hypothetical protein", Neurospora crassa	NC18F11.8	4e-04	17.7	13.8	17.7	21.1	Zn(II)Cys6
	Pc20g12860	strong similarity to transcription regulator of maltose utilization amyR - Aspergillus nidulans	gene: "104H10.220", product: "related to transcription activator amyR",	NC104H10.21	5e-63	5.2	8.4	6.0	6.8	Zn(II)Cys6
	Pc21g12520	weak similarity to transcription regulator of maltose utilization amyR - Aspergillus oryzae	transcription regulator SPBC1665.17 - fission yeast	T39609	2e-04	22.6	18.4	24.9	26.2	Zn(II)Cys6
	Pc22g07890	strong similarity to transcription regulator of maltose utilization amyR - Aspergillus oryzae	gene: "104H10.220", product: "related to transcription activator amyR",	NC104H10.21	2e-22	49.4	49.0	48.0	51.8	Zn(II)Cys6, Fungal specific transcription factor domain
	Pc22g12360	strong similarity to transcription regulator of maltose utilization amyR - Aspergillus oryzae	gene: "B23L1.140", product: "conserved hypothetical protein",	NCB23L1.20	0.0	121.5	83.0	333.7	458.8	Zn(II)Cys6, Fungal specific transcription factor domain
	Pc22g17220	weak similarity to transcription regulator of maltose utilization amyR - Aspergillus oryzae	gene: "B11AS.030", product: "conserved hypothetical protein",	NCB11AS.2	1e-37	34.8	32.5	31.3	44.2	Zn(II)Cys6, Fungal specific transcription factor domain
	An01g04370	strong similarity to transcription factor like protein An01g04370 - Aspergillus niger	zinc-finger protein - fission yeast (Schizosaccharomyces pombe)	T31762	6e-42	58.6	106.5	49.7	39.2	C2H2
	An01g04240	strong similarity to SWI SNF complex subunit like protein An01g04240 - Aspergillus niger	hypothetical protein SPCC285.17 - fission yeast	T41263	2e-17	122.2	123.3	84.7	74.4	DEK C-terminal domain, SWI60MDM2 domain
	An01g04600	strong similarity to gas transcription activator domain interacting protein like protein An01g04600 - Aspergillus niger	gene: "B23L1.140", product: "conserved hypothetical protein",	NCB23L1.20	0.0	14.0	11.1	11.1	11.6	BRCA1 C terminus (BRCT) domain
An02g06940	An02g02190	strong similarity to human regulatory molecule HRM-7 like protein An02g02190 - Aspergillus niger	product: "putative mitochondrial protein Tuba1p", Tuber borchii	AY27311.1	2e-52	266.9	275.8	226.2	276.6	No domain identified
	An02g05220	strong similarity to transcription regulator like protein An02g05220 - Aspergillus niger	product: "RING zinc finger protein SMR2", Homo sapiens striated	AF361946.1	3e-08	112.1	107.7	99.8	85.3	C3HC4 type (RING finger), zinc finger ZZ type, SH3 (Src homology 3) domain
	Pc13g04440	strong similarity to transcription factor like protein An02g06940 - Aspergillus niger	gene: "Tc1cp23", product: "Tc1cp23 protein", Mus musculus	BC055035.1	7e-15	5.3	7.5	7.1	5.9	CP2 transcription factor
	Pc16g06330	strong similarity to transcription factor like protein An02g06940 - Aspergillus niger [putative pseudogene]	product: "p61 MGR", Mus musculus p61 MGR mRNA, complete cds.	AF411211.1	1e-12	2.4	1.6	7.3	3.5	No domain identified
	Pc30g00420	strong similarity to transcription factor like protein An02g06940 - Aspergillus niger	product: "p61 MGR", Mus musculus p61 MGR mRNA, complete cds.	AF411211.1	1e-12	2.4	1.6	7.3	3.5	CP2 transcription factor
	Pc42g02200	strong similarity to transcription factor like protein An02g06940 - Aspergillus niger	hypothetical protein SPBC3084.04c - fission yeast	T40170	1e-39	328.0	299.2	259.0	225.1	ARID/BRH7 DNA binding domain
	An02g07140	strong similarity to transcription factor like protein An02g07140 - Aspergillus niger	probable histone acetyl transferase - fission yeast	T39004	1e-131	108.3	85.6	100.0	86.0	MOZ/SAS family
	An02g07140	strong similarity to transcription factor like protein An02g07140 - Aspergillus niger	gene: "CG11330", product: "SDO411370", Drosophila melanogaster	BT00941.1	4e-63	143.9	135.5	69.5	67.8	PHD-finger, JmC domain
	Pc22g09070	strong similarity to transcription factor like protein An02g07140 - Aspergillus niger	gene: "BTF18.040", product: "related to transcription factor TMF",	NCB718.3	3e-81	90.5	77.6	69.7	53.0	No domain identified
	An02g14310	strong similarity to PHD-finger protein like protein An02g14310 - Aspergillus niger	Gleobacter violaceus PCC 7421, complete genome, section	AP006574.135	3e-38	30.8	48.2	64.1	56.2	Pinn domain
An09g04970	An09g05800	strong similarity to transcription regulator like protein An09g05800 - Aspergillus niger	gene: "3H10.130", product: "related to regulator protein num1",	NC3H10.12	0.0	17.6	136.4	127.4	97.6	JmN domain, ARID/BRH7 DNA binding domain, PHD-finger, JmC domain
	An1g04480	strong similarity to non-specific RNA polymerase II transcription factor like protein An1g04480 - Aspergillus niger	transcription regulator SPBC1665.17 - fission yeast	T39609	2e-04	22.6	18.4	24.9	26.2	Zn(II)Cys6
	Pc21g11700	strong similarity to transcription activator like protein An21g00210 - Aspergillus niger	probable snf2 family helicase - fission yeast (Schizosaccharomyces pombe)	T37508	0.0	100.0	85.3	67.0	59.2	Res II (Type III restriction enzyme, res subunit), SNF2 family N-terminal
	Pc22g04070	strong similarity to human transcriptional regulator protein like protein An22g08420 - Aspergillus niger	gene: "smc4", product: "structural maintenance of chromosomes	AY081008.1	0.0	31.2	38.7	25.1	36.7	RecF/RecHSMC (structural maintenance of chromosomes) N terminal
	An12g10110	strong similarity to transcription factor like protein An12g10110 - Aspergillus niger	gene: "B16B8.160", product: "related to zinc finger protein crol	BX426314.16	3e-41	87.3	83.9	62.4	40.1	No domain identified
	An13g00370	strong similarity to acvA gene expression regulator Pcb04 like protein An13g00370 - Aspergillus niger	regulatory protein ARG2 - yeast (Saccharomyces cerevisiae)	S49827	2e-04	30.0	69.0	46.5	114.3	No domain identified
	An13g00370	strong similarity to acvA gene expression regulator Pcb04 like protein An13g00370 - Aspergillus niger	S. cerevisiae ARG1 regulatory gene	SCARGR1.1	4e-09	15.9	43.2	132.4	148.9	Zn(II)Cys6
	An13g00370	strong similarity to acvA gene expression regulator Pcb04 like protein An13g00370 - Aspergillus niger	probable pilbox factor - fission yeast (Schizosaccharomyces pombe)	T39995	2e-05	162.5	143.1	131.6	109.8	RNA recognition motif (RRM1 motif)
	An14g03170	strong similarity to collagenase zinc-finger transcription factor like protein An14g03170 - Aspergillus niger	gene: "14A2382.2", product: "14A2382.2 (novel protein similar to	MS42382.2	2e-38	33.1	41.4	30.6	23.2	CP2 transcription factor, No domain identified
	An14g04590	strong similarity to transcription factor like protein An14g04590 - Aspergillus niger	hypothetical protein, Dicyclothemium discoidium chromosome	AC117005.21	4e-11	101.2	96.3	68.7	57.1	Histone-like transcription factor (CBF/NFY)
An15g01640	Pc21g07940	strong similarity to transcription factor like protein An14g04590 - Aspergillus niger	unannotated ORF, Sequence 22 from Patent WO0059324.	NC115085.1	1e-79	17.8	14.2	85.8	66.3	PK (inositol polyphosphate kinase) domain
	Pc21g07940	strong similarity to transcription factor like protein An14g04590 - Aspergillus niger	gene: "H2CA", product: "H2CA protein", Emmericella nidulans	EMN24750.1	2e-04	23.7	21.9	20.4	20.4	No domain identified
	Pc22g19570	strong similarity to kvf gene expression regulator like protein An16g05120 - Aspergillus niger	gene: "pro1", product: "PRO1 protein", Sordaria brevicollis pro1 gene	SBR28536.1	1e-06	32.1	42.4	49.1	73.1	Zn(II)Cys6
	Pc21g11090	strong similarity to zinc-finger transcription factor like protein An16g08800 - Aspergillus niger	regulatory protein ARG2 - yeast (Saccharomyces cerevisiae)	S49827	2e-04	30.0	69.0	46.5	114.3	No domain identified
	Pc21g11090	strong similarity to zinc-finger transcription factor like protein An16g08800 - Aspergillus niger	S. cerevisiae ARG1 regulatory gene	SCARGR1.1	4e-09	15.9	43.2	132.4	148.9	Zn(II)Cys6
	An18g01300	strong similarity to transcription factor ap1p - Schizosaccharomyces pombe	similes: "B11ES.460", product: "putative protein", Neurospora crassa	BX46260.46	8e-06	20.8	16.4	12.8	12.8	Zn(II)Cys6
	Pc22g06330	similarity to transcription factor ap1p - Schizosaccharomyces pombe	regulatory protein ARG2 - yeast (Saccharomyces cerevisiae)	S49827	8e-05	43.1	44.8	60.6	85.0	Zn(II)Cys6
	Pc20g15780	similarity to transcription factor ap1p - Schizosaccharomyces pombe	regulatory protein ARG2 - yeast (Saccharomyces cerevisiae)	S49827	1e-13	12.5	18.7	31.4	55.3	Zn(II)Cys6
	Pc20g14370	similarity to transcription factor ap1p - Schizosaccharomyces pombe	gene: "VTE5.150", product: "conserved hypothetical protein",	NC17E5.14	8e-09	26.2	18.0	21.9	22.6	Zn(II)Cys6
	Pc20g10640	similarity to transcription factor ap1p - Schizosaccharomyces pombe	gene: "B3E4.120", product: "related to negative acting factor",	NCB3E4.9	8e-23	91.1	90.7	75.4	72.1	Zn(II)Cys6
AP1P	Pc22g08330	similarity to transcription factor ap1p - Schizosaccharomyces pombe	S. cerevisiae ARG1 regulatory gene	SCARGR1.1	4e-08	25.2	45.4	41.0	71.9	Zn(II)Cys6
	Pc22g10470	weak similarity to transcription factor ap1p - Schizosaccharomyces pombe	gene: "B9B15.005", product: "putative protein", Neurospora crassa	NCB9B15.1	2e-04	79.2	78.5	79.5	87.6	Zn(II)Cys6
	Pc22g12710	weak similarity to hypothetical transcription factor ap1p - Schizosaccharomyces pombe	gene: "B11ES.460", product: "putative protein", Neurospora crassa	BX46260.46	3e-06	91.5	88.7	38.7	36.1	Zn(II)Cys6
	Pc22g12710	weak similarity to hypothetical transcription factor ap1p - Schizosaccharomyces pombe	regulatory protein ARG2 - yeast (Saccharomyces cerevisiae)	S49827	2e-10	5.0	5.9	5.1	6.1	Zn(II)Cys6
	Pc22g19280	weak similarity to transcription factor ap1p - Schizosaccharomyces pombe	S. cerevisiae ARG1 regulatory gene	SCARGR1.1	4e-05	8.5	17.3	12.0	12.0	Zn(II)Cys6
	Pc22g19540	weak similarity to transcription factor ap1p - Schizosaccharomyces pombe	gene: "B3E4.120", product: "related to negative acting factor",	NCB3E4.9	3e-10	193.2	157.1	113.0	86.6	Zn(II)Cys6
	Pc22g23360	similarity to transcription factor ap1p - Schizosaccharomyces pombe	gene: "BTF21.070", product: "hypothetical protein", Neurospora crassa	NCB7F21.6	7e-07	25.4	35.7	36.9	45.5	Zn(II)Cys6
	Pc22g06330	strong similarity to regulator protein of the arginine metabolism ARG2 - Saccharomyces cerevisiae	regulatory protein ARG2 - yeast (Saccharomyces cerevisiae)	S49827	2e-04	30.0	69.0	46.5	114.3	No domain identified
	Pc12g02910	weak similarity to transcription factor ap1p - Schizosaccharomyces pombe	regulatory protein ARG2 - yeast (Saccharomyces cerevisiae)	S49827	2e-04	30.0	69.0	46.5	114.3	No domain identified
	Pc12g13730	weak similarity to transcription factor ap1p - Schizosaccharomyces pombe	regulatory protein ARG2 - yeast (Saccharomyces cerevisiae)	S49827	2e-04	30.0	69.0	46.5	114.3	No domain identified
ARCA	Pc30g03330	similarity to transcription factor ap1p - Schizosaccharomyces pombe	S. cerevisiae ARG1 regulatory gene	SCARGR1.1	4e-09	15.9	43.2	132.4	148.9	Zn(II)Cys6
	Pc13g07020	similarity to regulator protein ap1p - Schizosaccharomyces pombe	regulatory protein ARG2 - yeast (Saccharomyces cerevisiae)	S49827	2e-04	30.0	69.0	46.5	114.3	No domain identified
	Pc16g04470	similarity to transcription factor ap1p - Schizosaccharomyces pombe	gene: "pro1", product: "PRO1 protein", Sordaria macrospora pro1	SAJ1051.1	8e-06	148.1	133.2	118.7	124.4	Zn(II)Cys6
	Pc16g07620	similarity to hypothetical transcription factor ap1p - Schizosaccharomyces pombe	similes: "B11ES.460", product: "putative protein", Neurospora crassa	BX46260.46	8e-06	20.8	16.4	12.8	12.8	Zn(II)Cys6
	Pc16g10840	similarity to transcription factor ap1p - Schizosaccharomyces pombe	regulatory protein ARG2 - yeast (Saccharomyces cerevisiae)	S49827	8e-05	43.1	44.8	60.6	85.0	Zn(II)Cys6
	Pc20g04200	similarity to transcription factor ap1p - Schizosaccharomyces pombe	regulatory protein ARG2 - yeast (Saccharomyces cerevisiae)	S49827	1e-13	12.5	18.7	31.4	55.3	Zn(II)Cys6
	Pc20g06630	similarity to transcription factor ap1p - Schizosaccharomyces pombe	gene: "VTE5.150", product: "conserved hypothetical protein",	NC17E5.14	8e-09	26.2	18.0	21.9	22.6	Zn(II)Cys6
	Pc20g08330	weak similarity to transcription factor ap1p - Schizosaccharomyces pombe	gene: "B3E4.120", product: "related to negative acting factor",	NCB3E4.9	8e-23	91.1	90.7	75.4	72.1	Zn(II)Cys6
	Pc20g15780	similarity to transcription factor ap1p - Schizosaccharomyces pombe	S. cerevisiae ARG1 regulatory gene	SCARGR1.1	4e-08	25.2	45.4	41.0	71.9	Zn(II)Cys6
	Pc20g14370	similarity to transcription factor ap1p - Schizosaccharomyces pombe	gene: "B9B15.005", product: "putative protein", Neurospora crassa	NCB9B15.1	2e-04	79.2	78.5	79.5	87.6	Zn(II)Cys6
AR081	Pc22g10470	weak similarity to transcription factor ap1p - Schizosaccharomyces pombe	gene: "B11ES.460", product: "putative protein", Neurospora crassa	BX46260.46	3e-06	91.5	88.7	38.7	36.1	Zn(II)Cys6
	Pc22g12710	weak similarity to hypothetical transcription factor ap1p - Schizosaccharomyces pombe	regulatory protein ARG2 - yeast (Saccharomyces cerevisiae)	S49827	2e-10	5.0	5.9	5.1	6.1	Zn(II)Cys6
	Pc22g19280	weak similarity to transcription factor ap1p - Schizosaccharomyces pombe	S. cerevisiae ARG1 regulatory gene	SCARGR1.1	4e-05	8.5	17.3	12.0	12.0	Zn(II)Cys6
	Pc22g19540	weak similarity to transcription factor ap1p - Schizosaccharomyces pombe	gene: "B3E4.120", product: "related to negative acting factor",	NCB3E4.9	3e-10	193.2	157.1	113.0	86.6	Zn(II)Cys6
	Pc22g23360	similarity to transcription factor ap1p - Schizosaccharomyces pombe	gene: "BTF21.070", product: "hypothetical protein", Neurospora crassa	NCB7F21.6	7e-07	25.4	35.7	36.9	45.5	Zn(II)Cys6
	Pc22g06330	strong similarity to regulator protein of the arginine metabolism ARG2 - Saccharomyces cerevisiae	regulatory protein ARG2 - yeast (Saccharomyces cerevisiae)	S49827	2e-04	30.0	69.0	46.5	114.3	No domain identified
	Pc12g02910	weak similarity to transcription factor ap1p - Schizosaccharomyces pombe	regulatory protein ARG2 - yeast (Saccharomyces cerevisiae)	S49827	2e-04	30.0	69.0	46.5	114.3	No domain identified
	Pc12g13730	weak similarity to transcription factor ap1p - Schizosaccharomyces pombe	regulatory protein ARG2 - yeast (Saccharomyces cerevisiae)	S49827	2e-04	30.0	69.0	46.5	114.3	No domain identified
	Pc30g03330	similarity to transcription factor ap1p - Schizosaccharomyces pombe	S. cerevisiae ARG1 regulatory gene	SCARGR1.1	4e-09	15.9	43.2	132.4	148.9	Zn(II)Cys6
	Pc13g07020	similarity to regulator protein ap1p - Schizosaccharomyces pombe	regulatory protein ARG2 - yeast (Saccharomyces cerevisiae)	S49827	2e-04	30.0	69.0	46.5	114.3	No domain identified
AR082	Pc16g04470	similarity to transcription factor ap1p - Schizosaccharomyces pombe	gene: "pro1", product: "PRO1 protein", Sordaria macrospora pro1	SAJ1051.1	8e-06	148.1	133.2	118.7	124.4	Zn(II)Cys6
	Pc16g07620	similarity to hypothetical transcription factor ap1p - Schizosaccharomyces pombe	similes: "B11ES.460", product: "putative protein", Neurospora crassa	BX46260.46	8e-06	20.8	16.4	12.8	12.8	Zn(II)Cys6
	Pc16g10840	similarity to transcription factor ap1p - Schizosaccharomyces pombe	regulatory protein ARG2 - yeast (Saccharomyces cerevisiae)	S49827	8e-05	43.1	44.8	60.6	85.0	Zn(II)Cys6
	Pc20g04200	similarity to transcription factor ap1p - Schizosaccharomyces pombe	regulatory protein ARG2 - yeast (Saccharomyces cerevisiae)	S49827	1e-13	12.5	18.7	31.4	55.3	Zn(II)Cys6
	Pc20g06630	similarity to transcription factor ap1p - Schizosaccharomyces pombe	gene: "VTE5.150", product: "conserved hypothetical protein",	NC17E5.14	8e-09	26.2	18.0	21.9	22.6	Zn(II)Cys6
	Pc20g08330	weak similarity to transcription factor ap1p - Schizosaccharomyces pombe	gene: "B3E4.120", product: "related to negative acting factor",	NCB3E4.9	8e-23	91.1	90.7	75.4	72.1	Zn(II)Cys6
	Pc20g15780	similarity to transcription factor ap1p - Schizosaccharomyces pombe	S. cerevisiae ARG1 regulatory gene	SCARGR1.1	4e-08					

Hypothetical transcription regulator	Pc22at1860	strong similarity to hypothetical transcription factor - Schizosaccharomyces pombe	probable transcription factor - fission yeast (Schizosaccharomyces pombe)	T37869	4e-40	74.2	59.0	43.0	49.4	YEATS family
	Pc14g00020	weak similarity to hypothetical transcription regulator protein - Schizosaccharomyces pombe	gene: "SE6.310": product: "related to nitrate assimilation regulatory protein - fission yeast"	NC568_27	4e-24	85.6	122.7	163.2	348.8	Zn(II)Cys6
	Pc16t1780	weak similarity to hypothetical transcription regulator protein - Schizosaccharomyces pombe	gene: "SPBC250.11c": product: "fission yeast"	T40327	2e-05	55.4	55.7	49.5	75.4	Zn(II)Cys6 Fungal specific transcription factor domain
	Pc18g1980	strong similarity to hypothetical transcription regulator protein - Schizosaccharomyces pombe	gene: "SPAC1327.01c": "SPAC1836.16c": "SPAC1783.09c":	SPAC1327_1	4e-60	81.3	89.0	71.9	78.1	Zn(II)Cys6 Fungal specific transcription factor domain
	Pc19g0420	weak similarity to hypothetical transcription regulator protein - Schizosaccharomyces pombe	gene: "SPAPB1A11.04c": "SPAC6 chromosomes I BAC pb1A11.1"	SPAPB1A11_4	5e-15	20.5	23.6	38.7	77.8	Zn(II)Cys6 Fungal specific transcription factor domain
	Pc20g0620	weak similarity to hypothetical transcription regulator protein - Schizosaccharomyces pombe	gene: "SPAC11.04c": product: "fission yeast"	SPAC11_4	4e-10	34.5	41.0	26.4	35.1	Zn(II)Cys6 Fungal specific transcription factor domain
	Pc21g0230	weak similarity to hypothetical transcription regulator protein - Schizosaccharomyces pombe	hypothetical protein SPBC530.11c - fission yeast	T40527	6e-11	25.9	19.6	22.9	31.4	Fungal specific transcription factor domain
	Pc22g0620	weak similarity to hypothetical transcription regulator protein - Schizosaccharomyces pombe	gene: "B15B3.150": product: "related to general repressor of"	BX84262_15	2e-14	41.3	23.9	65.3	31.9	Fungal specific transcription factor domain
	Pc22at1880	weak similarity to hypothetical transcription regulator protein - Schizosaccharomyces pombe	gene: "SPAPB1A11.04c": S. pombe chromosome I BAC pb1A11.1"	SPAPB1A11_4	7e-09	21.9	28.4	34.4	60.3	Zn(II)Cys6 Fungal specific transcription factor domain
	Pc21g1270	weak similarity to growth regulator BKT - Mus musculus	hypothetical protein SPAC45.18c - Mus musculus	SPAC45_18	5e-15	59.4	54.3	54.4	61.2	PHD finger
ING1	Pc21g1570	similarity to JUN-like transcription factor JbA - Aspergillus nidulans	gene: "JbA": product: "JUN-like b2P transcription factor"; Emicella	JB481222_1	4e-11	17.2	17.4	13.0	16.5	b2P 1 and b2P 2 (Basic region leucine zipper)
JLBA	Pc16g6680	similarity to positive regulator of the lactose-galactose induction LAC9 - Kluyveromyces lactis	gene: "B2E7.100": product: "related to lactose regulatory protein";	BX697675_1	1e-103	253.6	257.1	275.1	336.7	Zn(II)Cys6 Fungal specific transcription factor domain
LAC9	Pc20g3300	similarity to positive regulator of the lactose-galactose induction LAC9 - Kluyveromyces lactis	product: "Unknown protein"; Anabaena thaliana Unknown protein	BT01680_1	0-15	150.0	125.8	143.6	118.4	Zn(II)Cys6 Fungal specific transcription factor domain
LaE	Pc16t14010	strong similarity to hypothetical methyltransferase ACO3467.1 - Oberea zea	Aspergillus fumigatus LaEa gene, complete cds	AF127723_1	1e-130	355.6	361.3	374.8	338.3	Methyltransferase domain
LEU3	Pc12g00620	similarity to leucine-specific regulatory protein Leu3 - Saccharomyces cerevisiae	regulatory protein LEU3 - yeast (Saccharomyces cerevisiae)	RB0Y13	1e-14	26.8	19.0	29.0	20.4	Zn(II)Cys6
	Pc20g07510	strong similarity to leucine-specific regulatory protein Leu3 - Saccharomyces cerevisiae	regulatory protein LEU3 - yeast (Saccharomyces cerevisiae)	RB0Y13	8e-44	238.5	211.1	174.3	194.5	Zn(II)Cys6 Fungal specific transcription factor domain
	Pc06g01850	weak similarity to transcription activator of lysine pathway Lyt4 - Saccharomyces cerevisiae	hypothetical protein YGR388w - yeast (Saccharomyces cerevisiae)	S84623	10-1014	177.6	117.2	119.5	124.7	Zn(II)Cys6
	Pc12g01430	similarity to transcription activator of lysine pathway Lyt4 - Saccharomyces cerevisiae [putative pseudogene]	DNA centromere region sequence from BAC DP26806, DP34F04, gene: "17E5.150": product: "conserved hypothetical protein";	CNS09S45_32	1e-31	27.1	15.0	31.7	32.7	Zn(II)Cys6
	Pc12g0570	weak similarity to transcription activator of lysine pathway Lyt4 - Saccharomyces cerevisiae	product: "putative cysteine-binding protein (bP)"; Histella volcani	NC17E5_14	7e-09	2.2	2.8	4.1	2.3	Zn(II)Cys6
	Pc13g0220	similarity to transcription activator of lysine pathway Lyt4 - Saccharomyces cerevisiae	hypothetical fungal Ziv2-Cys6(z) zinc-finger protein - fission yeast	AF15825_1	6e-58	94.5	92.5	59.5	61.7	Zn(II)Cys6
	Pc13g0580	similarity to transcription activator of lysine pathway Lyt4 - Saccharomyces cerevisiae	gene: "MPP1": product: "Zn(II)Cys6 transcription factor"; Pichia	T41718	9e-06	89.4	50.3	42.5	53.7	Zn(II)Cys6
	Pc14g00980	similarity to transcription activator of lysine pathway Lyt4 - Saccharomyces cerevisiae	hypothetical fungal Ziv2-Cys6(z) zinc-finger protein - fission yeast	AY100521_1	2e-16	14.0	8.4	11.8	16.4	Zn(II)Cys6
	Pc20g1750	similarity to transcription activator of lysine pathway Lyt4 - Saccharomyces cerevisiae	transcription activator LYt4 - yeast (Saccharomyces cerevisiae)	T41718	1e-06	44.2	69.4	153.7	162.8	Zn(II)Cys6
	Pc20g02030	similarity to transcription activator of lysine pathway Lyt4 - Saccharomyces cerevisiae	probable membrane protein YDR213w - yeast (Saccharomyces cerevisiae)	S81937	3e-04	24.3	18.5	37.9	39.3	Zn(II)Cys6
	Pc20g10800	similarity to transcription activator of lysine pathway Lyt4 - Saccharomyces cerevisiae	gene: "Ca2OC1.12": product: "possible zinc-finger protein"; Caldicoccus	S61580	2e-06	73.0	60.1	99.5	138.1	Zn(II)Cys6
	Pc20g15410	weak similarity to transcription activator of lysine pathway Lyt4 - Saccharomyces cerevisiae	transcription activator LYt4 - yeast (Saccharomyces cerevisiae)	CAC2OC1_11	2e-04	6.5	6.2	11.8	13.7	Zn(II)Cys6
	Pc19g00920	similarity to transcription activator of lysine pathway Lyt4 - Saccharomyces cerevisiae	gene: "meaA": product: "Yeast (Saccharomyces cerevisiae) MeaA protein"	S61587	8e-05	44.6	37.0	40.2	46.2	Zn(II)Cys6
	Pc19g05540	weak similarity to transcription activator of lysine pathway Lyt4 - Saccharomyces cerevisiae	gene: "meaB": product: "developmental regulator meB"; Aspergillus	BN16862_1	1e-104	169.6	125.2	97.8	101.5	Zn(II)Cys6
	Pc19g07600	weak similarity to transcription activator of lysine pathway Lyt4 - Saccharomyces cerevisiae	gene: "ARCA": product: "ARCA protein"; Emicella nidulans ARCA	EN204750_1	1e-116	169.6	125.2	97.8	101.5	Zn(II)Cys6
	Pc22g02680	weak similarity to transcription activator of lysine pathway Lyt4 - Saccharomyces cerevisiae	gene: "SE6.190": product: "related to ARCA protein"; Neurospora	NC556_17	1e-15	48.0	60.8	24.8	23.6	Zn(II)Cys6
	Pc22g04260	weak similarity to transcription activator of lysine pathway Lyt4 - Saccharomyces cerevisiae	product: "hypothetical protein"; Aspergillus fumigatus BAC plct project	BX64605_122	1e-62	68.8	61.7	58.5	68.6	Zn(II)Cys6
	Pc22g04500	weak similarity to transcription activator of lysine pathway Lyt4 - Saccharomyces cerevisiae	gene: "AA12H2.19c": product: "hypothetical protein"; Aspergillus	AA12H2_19	2e-64	2.9	2.9	15.2	11.6	Zn(II)Cys6 Fungal specific transcription factor domain
	Pc13g02490	weak similarity to maltose pathway regulatory protein Mal13 - Saccharomyces cerevisiae	gene: "B1014.250": product: "hypothetical protein"; Neurospora	NCB1014_25	2e-14	0.3	0.5	0.7	0.4	Zn(II)Cys6
	Pc16g08120	similarity to maltose pathway regulatory protein Mal13 - Saccharomyces cerevisiae	hypothetical protein YGR388w - yeast (Saccharomyces cerevisiae)	S84623	4e-08	10.1	10.4	11.7	11.4	Zn(II)Cys6
	Pc19g07200	similarity to maltose pathway regulatory protein Mal13 - Saccharomyces cerevisiae	gene: "104H10.220": product: "related to transcription activator amYr";	NC104H10_21	3e-11	69.0	171.3	63.1	75.1	Zn(II)Cys6
	Pc20g15410	strong similarity to MADS-box domain transcription factor Momi - Saccharomyces cerevisiae	gene: "AA12H2.19c": product: "hypothetical protein"; Aspergillus	AA12H2_19	3e-07	209.2	192.3	161.7	134.7	Zn(II)Cys6
MCMI	Pc19g02880	strong similarity to MADS-box domain transcription factor Momi - Saccharomyces cerevisiae	Yeast (Saccharomyces cerevisiae) FUMO gene, complete cds	AF15825_1	5e-58	94.5	92.5	59.5	61.7	MADS-box/SRF-type transcription factor (DNA-binding and dimerisation)
MEAB	Pc19g02880	strong similarity to MADS-box domain transcription factor Momi - Saccharomyces cerevisiae	gene: "meaB": product: "MEAB protein"; Aspergillus nidulans meaB	ANMEABG_1	1e-104	382.5	368.9	303.4	353.0	12P leucine zipper
MEDB	Pc20g08130	similarity to transcription factor MedB - Saccharomyces cerevisiae	hypothetical protein SPBC21.04 - fission yeast (Schizosaccharomyces pombe)	T39866	1e-12	153.5	117.8	166.9	106.6	No domain identified
MEDA (MEDUSA)	Pc22g13450	strong similarity to hypothetical Medusa medA - Aspergillus nidulans	gene: "meaA": product: "Medusa"; Emicella nidulans Medusa	AF080599_1	7e-102	13.5	14.4	85.7	52.7	No domain identified
	Pc22g13450	weak similarity to developmental regulator medA - Aspergillus nidulans	gene: "meaB": product: "developmental regulator medA"; Aspergillus	AF207933_1	1e-104	155.2	140.9	101.5	101.5	No domain identified
MPP1	Pc12g02890	similarity to hypothetical Zn(II)Cys6 transcription factor MPP1 - Pichia angusta	gene: "MPP1": product: "Zn(II)Cys6 transcription factor"; Pichia	AY100521_1	4e-20	31.0	27.8	26.1	23.3	No domain identified
MTFB	Pc14g02020	strong similarity to hypothetical Zn(II)Cys6 transcription factor MPP1 - Pichia angusta	gene: "MPP1": product: "Zn(II)Cys6 transcription factor"; Pichia	AY100521_1	1e-36	21.5	23.9	27.1	34.1	No domain identified
NC2	Pc12g03700	weak similarity to mitochondrial transcription factor mTFB - Kluyveromyces lactis	gene: "NC2": product: "NC2 protein"; Kluyveromyces lactis	SPAC1002.08c	1e-14	42.5	37.5	38.5	42.5	HAAD domain (ribosomal RNA adenine dimethylase)
NF-X1	Pc12g13040	similarity to alpha chain of transcription factor NF-X1 - Homo sapiens	probable transcription repressor - fission yeast	T37869	3e-18	158.5	139.0	146.4	96.5	Core histone H2AF(2B)H3H4. Histone-like transcription factor (CBF/NFY)
NIRA	Pc19g09140	strong similarity to cysteine-rich transcription factor NF-X1 - Homo sapiens	probable cysteine-rich transcription regulator - fission yeast	T41146	1e-144	41.8	43.6	34.7	36.5	NF-X1 type zinc finger, R3H domain
	Pc13g02000	weak similarity to nitrate assimilation regulatory protein nira - Aspergillus nidulans	nitrate assimilation regulatory protein nira - Emicella nidulans	AF1687	6e-37	19.7	18.3	20.0	24.3	Zn(II)Cys6 Fungal specific transcription factor domain
	Pc13g02410	weak similarity to nitrate assimilation regulatory protein nira - Aspergillus nidulans	gene: "SPBC1683.13c": Schizosaccharomyces pombe cosmid	SPBC1683_13	3e-32	41.8	43.6	52.8	41.6	Fungal specific transcription factor domain
	Pc13g02410	strong similarity to nitrate assimilation regulatory protein nira - Aspergillus nidulans	nitrate assimilation regulatory protein nira - Emicella nidulans	AF1687	0-10	105.1	75.8	57.6	55.5	Zn(II)Cys6 Fungal specific transcription factor domain
	Pc06g00120	weak similarity to nitrogen assimilation regulatory protein nit-4 - Neurospora crassa	related to pathway-specific nitrogen regulator [imported] - Neurospora	T51029	1e-137	428.4	335.3	112.8	100.3	Zn(II)Cys6 Fungal specific transcription factor domain
	Pc12g00270	strong similarity to nitrogen assimilation regulatory protein nit-4 - Neurospora crassa	regulatory protein nit-4 - Neurospora crassa	AF1696	4e-30	70.5	86.5	112.1	170.4	Zn(II)Cys6 Fungal specific transcription factor domain
	Pc12g09830	weak similarity to nitrogen assimilation regulatory protein nit-4 - Neurospora crassa [putative sequencing error]	regulatory protein nit-4 - Neurospora crassa	AF1696	2e-37	179.5	161.9	133.2	107.8	Zn(II)Cys6 Fungal specific transcription factor domain
	Pc12g11450	similarity to nitrogen assimilation regulatory protein nit-4 - Neurospora crassa	regulatory protein nit-4 - Neurospora crassa	AF1696	4e-08	299.2	258.7	332.3	283.6	Zn(II)Cys6
	Pc13g15600	weak similarity to nitrogen assimilation regulatory protein nit-4 - Neurospora crassa	gene: "MPP1": product: "Zn(II)Cys6 transcription factor"; Pichia	AY100521_1	3e-04	64.5	61.1	36.7	37.5	Zn(II)Cys6
	Pc16g09100	weak similarity to nitrogen assimilation regulatory protein nit-4 - Neurospora crassa	regulatory protein nit-4 - Neurospora crassa	AF1696	2e-37	179.5	161.9	133.2	107.8	Fungal specific transcription factor domain
	Pc16g10210	weak similarity to nitrogen assimilation regulatory protein nit-4 - Neurospora crassa	hypothetical protein SPBC530.05 - fission yeast	T40521	2e-06	48.5	39.4	58.3	62.9	Zn(II)Cys6
	Pc16g13230	similarity to nitrogen assimilation regulatory protein nit-4 - Neurospora crassa	regulatory protein nit-4 - Neurospora crassa	AF1696	2e-35	23.1	17.7	16.5	11.0	Zn(II)Cys6 Fungal specific transcription factor domain
	Pc21g12800	strong similarity to nitrogen assimilation regulatory protein nit-4 - Neurospora crassa	regulatory protein nit-4 - Neurospora crassa	AF1696	9e-45	0.5	0.8	1.5	2.2	Zn(II)Cys6
	Pc22g06340	weak similarity to nitrogen assimilation regulatory protein nit-4 - Neurospora crassa	gene: "SPAPB1A11.04c": S. pombe chromosome I BAC pb1A11.1"	SPAPB1A11_4	7e-09	21.9	28.4	34.4	60.3	Zn(II)Cys6 Fungal specific transcription factor domain
	Pc22g14440	similarity to nitrogen assimilation regulatory protein nit-4 - Neurospora crassa	gene: "SPBC1683.13c": Schizosaccharomyces pombe cosmid	SPBC1683_13	3e-31	18.5	23.1	20.6	32.6	Zn(II)Cys6 Fungal specific transcription factor domain
	Pc22g14550	weak similarity to nitrogen assimilation regulatory protein nit-4 - Neurospora crassa	gene: "X4G11.050": product: "hypothetical protein"; Neurospora crassa	NCX4G11_5	2e-04	129.4	186.6	125.2	136.4	Fungal specific transcription factor domain
	Pc22g0620	strong similarity to nitrogen assimilation regulatory protein nit-4 - Neurospora crassa	regulatory protein nit-4 - Neurospora crassa	AF1696	2e-37	179.5	161.9	133.2	107.8	Zn(II)Cys6 Fungal specific transcription factor domain
NOT3	Pc20g07490	strong similarity to transcription regulator not3 - Homo sapiens	probable transcription regulator - fission yeast (Schizosaccharomyces pombe)	T37869	3e-18	158.5	139.0	146.4	96.5	Core histone H2AF(2B)H3H4. Histone-like transcription factor (CBF/NFY)
	Pc22g14600	GATA transcription factor nr6B - Penicillium chrysogenum	gene: "nr6B": product: "GATA transcription factor"; Penicillium	U9385_1	1e-146	76.4	78.6	68.1	43.3	GATA zinc finger
	Pc22g16880	regulator of nitrogen metabolism repression nre - Penicillium chrysogenum [putative sequencing error]	NITROGEN REGULATORY PROTEIN AREA (NITROGEN)	AREA_PENRO	0-10	162.5	149.3	144.4	113.1	Nitrogen regulatory protein AreA N terminus, GATA zinc finger
	Pc12g05460	strong similarity to transcription activator nuc-1 - Neurospora crassa	probable transcription activator nuc-1 - Neurospora crassa	A38373	3e-40	108.2	65.5	90.3	70.7	Helix-loop-helix DNA-binding domain
	Pc23g00570	similarity to transcription activator nuc-1 - Neurospora crassa [truncated ORF] [putative sequencing error]	probable transcription activator nuc-1 - Neurospora crassa	A38373	5e-23	7.7	6.3	4.8	2.9	Helix-loop-helix DNA-binding domain
NUC-1	Pc18g04040	weak similarity to hypothetical transcription regulator PAB1642 - Pyrococcus abyssi	gene: "PF1337": product: "transcriptional activator, putative";	AE010238_9	2e-11	58.5	36.7	33.3	33.5	TENAT4-APQC family
PACC	Pc18g04040	transcription factor pacc - Penicillium chrysogenum	gene: "pacc": product: "transcription factor Pacc"; Penicillium	PCU44726_1	0-0	211.5	171.5	219.2	151.2	C2H2
PAB	Pc13g01950	similarity to transcription regulator 2 of paa operon pab - Bacillus subtilis	gene: "mlb013": product: "transcriptional regulator"; Mesorhizobium	AP00294_10	3e-30	4.8	3.4	7.1	5.5	FMN-binding domain 2
	Pc15g01210	similarity to transcription regulator 2 of paa operon pab - Bacillus subtilis	gene: "mlb013": product: "transcriptional regulator"; Mesorhizobium	AP00294_10	2e-36	88.3	83.5	174.2	169.7	FMN-binding domain 2
PBP	Pc13g15350	similarity to transcription factor PBP - Fusarium solani	gene: "nsd0": product: "Nsd0"; Aspergillus nidulans Nsd0 (nsd)	ENJ70043_1	1e-112	110.1	82.8	82.5	49.4	GATA zinc finger
	Pc22g02540	strong similarity to transcription factor PBP - Fusarium solani	gene: "nsd0": product: "Nsd0"; Aspergillus nidulans Nsd0 (nsd)	AF026072_2	3e-35	49.1	51.4	70.2	73.2	GATA zinc finger, PAS fold domain
PHD	Pc18g00330	strong similarity to hypothetical phd finger transcription regulator - Schizosaccharomyces pombe	probable phd finger transcription regulator - fission yeast	T41146	2e-32	36.0	29.3	27.4	27.3	PHD finger
	Pc12g13120	similarity to the transcription factor Pp2 - Saccharomyces cerevisiae	transcription factor PP2 - yeast (Saccharomyces cerevisiae)	S70646	2e-05	43.0	70.2	66.1	127.8	Zn(II)Cys6
	Pc13g00900	weak similarity to the transcription factor Pp2 - Saccharomyces cerevisiae	product: "transcriptional regulator, putative"; Aspergillus fumigatus	BX64607_85	5e-04	3.3	7.2	5.5	9.4	Zn(II)Cys6
	Pc20g1150	weak similarity to the transcription factor Pp2 - Saccharomyces cerevisiae	gene: "B9B11.100": product: "related to grin"; Neurospora crassa DNA	NCB9B11_10	1e-132	101.7	101.4	130.7	132.7	Prim. Trin C-terminal cupin domain, Cupin domain
PIRIN: Cofactor of	Pc06g02200	strong similarity to phosphorus acquisition transcriptional activator NUC-1 negative regulator PREG - Neurospora crassa	regulatory protein preg - Neurospora crassa	SC5274	1e-45	4.4	3.2	3.8	2.4	Cyclin domain
	Pc13g00960	strong similarity to phosphorus acquisition transcriptional activator NUC-1 negative regulator PREG - Neurospora crassa	regulatory protein preg - Neurospora crassa	SC5274	2e-46	13.2	8.3	7.1	4.1	Cyclin domain
	Pc20g1150	strong similarity to phosphorus acquisition transcriptional activator NUC-1 negative regulator PREG - Neurospora crassa	regulatory protein preg - Neurospora crassa	SC5274	1e-49	129.0	99.3	131.3	138.2	Cyclin domain, Cyclin N-terminal domain
	Pc12g11430	similarity to regulator protein Pp1 - Saccharomyces cerevisiae	gene: "B92.010": product: "conserved hypothetical protein";	NC896_1	4e-62	10.0	10.2	18.6	23.6	Zn(II)Cys6 Fungal specific transcription factor domain
	Pc12g14500	similarity to regulator protein Pp1 - Saccharomyces cerevisiae	probable membrane protein YHR78w - yeast (Saccharomyces cerevisiae)	S50366	3e-41	2.5	2.8	2.6	10.8	Zn(II)Cys6 Fungal specific transcription factor domain
	Pc16g07300	similarity to regulator protein Pp1 - Saccharomyces cerevisiae	probable membrane protein YHR78w - yeast (Saccharomyces cerevisiae)	S48404_1	2e-17	43.6	31.7	44.4	31.0	Zn(II)Cys6 Fungal specific transcription factor domain
	Pc20g15510	similarity to regulator protein Pp1 - Saccharomyces cerevisiae	probable regulatory protein YHR78w - yeast (Saccharomyces cerevisiae)	SA0917	5e-25	77.2	92.2	78.9	107.1	Zn(II)Cys6 Fungal specific transcription factor domain
	Pc22g04590	similarity to regulator protein Pp1 - Saccharomyces cerevisiae	regulatory protein PPR1 - yeast (Saccharomyces cerevisiae)	RB0Y11	3e-16	238.9	219.2			

QUTA	Pc13g08420	strong similarity to regulator protein qutA - Aspergillus nidulans	gene: "qutA"; product: "quinic acid utilization activator, putative";	BX49605_38	0.0	51.2	44.0	45.1	49.8	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
	Pc16g00390	strong similarity to regulator protein qutA - Aspergillus nidulans	transcription regulator, binuclear cluster zinc-finger protein - fission	T39677	2e-04	134.5	125.0	107.4	107.8	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
QUTR	Pc12g14140	similarity to regulator protein qutA - Aspergillus nidulans	gene: "MPPI1" product: "Zn(II)(Cys) ₆ transcription factor"; Pchla	AY100521_1	2e-05	36.6	54.4	61.3	69.9	Zn(II)(Cys) ₆	
	Pc22g12620	weak similarity to regulator protein qutA - Aspergillus nidulans	gene: "MPPI1" product: "Zn(II)(Cys) ₆ transcription factor"; Pchla	AY100521_1	1e-10	12.4	9.8	15.7	16.4	Zn(II)(Cys) ₆	
RDR1	Pc12g16820	strong similarity to repressor protein qutA - Aspergillus nidulans	gene: "MPPI1" product: "Zn(II)(Cys) ₆ transcription factor"; Pchla	AY100521_1	1e-10	12.4	9.8	15.7	16.4	Zn(II)(Cys) ₆	
	Pc12g02800	similarity to zinc-finger transcription factor Rdr1 - Saccharomyces cerevisiae	probable membrane protein YOR380w - yeast (Saccharomyces)	S67292	2e-85	221.1	222.7	224.9	193.7	Zn(II)(Cys) ₆	Uncharacterised protein family UPF0005
RES2p	Pc12g16580	strong similarity to zinc-finger transcription factor Rdr1 - Saccharomyces cerevisiae	probable membrane protein YOR380w - yeast (Saccharomyces)	S67292	7e-47	12.8	9.3	11.7	13.1	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
	Pc16g07940	weak similarity to transcription repressor Rdr1 - Saccharomyces cerevisiae	probable membrane protein YOR380w - yeast (Saccharomyces)	S67292	3e-45	13.0	12.1	11.0	11.3	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
RFX	Pc12g17990	strong similarity to zinc-finger transcription repressor Rdr1 - Saccharomyces cerevisiae	probable membrane protein YOR380w - yeast (Saccharomyces)	S67292	3e-45	13.0	12.1	11.0	11.3	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
	Pc12g07510	weak similarity to cell cycle regulator res2p - Schizosaccharomyces pombe	CELL DIVISION CYCLE RELATED-PROTEIN RES2PCT1	RES2_SCHPO	3e-62	108.6	89.8	73.2	69.8	APES domain	Arkylin repeat
RFXH	Pc21g03670	transcription factor rfxH - Penicillium chrysogenum	gene: "rfxH"; product: "RfxH"; Penicillium chrysogenum RfxH (rfxH)	AY189831_1	0.0	85.9	103.4	67.4	131.5	GATA zinc finger	
	Pc22g06930	similarity to transcription regulator Rfx1 - Candida albicans	gene: "Rfx1"; product: "hypothetical protein"; Neurospora crassa	NCB1821_14	2e-27	21.0	15.5	7.4	5.8	HMG (high mobility group) box	
RFX1	Pc20g16180	transcription factor like protein RFX - Penicillium chrysogenum	gene: "RFX1"; product: "putative RFX transcription factor"; Penicillium	PC243306_1	0.0	180.0	153.5	94.4	92.0	RFX DNA-binding domain	
	Pc21g11300	similarity to regulator protein Rfx1 - Saccharomyces cerevisiae	hypothetical protein SPBC104.10c: SPBP23A10.01c - fission yeast	T50388	2e-28	240.8	222.8	218.2	157.4	Mediator complex subunit MED14	
RFX2	Pc13g12780	strong similarity to regulator protein Rfx1 - Saccharomyces cerevisiae	gene: "B12N10.070"; product: "related to regulator of DNA centromeric region sequence from GALC-DP26806; DP34F04	NCB12N19_6	0.0	9.3	6.5	6.8	5.3	JMN domain, JMN1 domain	
	Pc22g06240	strong similarity to protein Rfx1 - Saccharomyces cerevisiae	gene: "B12N10.070"; product: "related to regulator of DNA centromeric region sequence from GALC-DP26806; DP34F04	NCB12N19_6	0.0	9.3	6.5	6.8	5.3	JMN domain, JMN1 domain	
RFX3	Pc22g06620	strong similarity to chromatin remodeling gene transcription regulator Rfx2 - Saccharomyces cerevisiae	gene: "B12N10.070"; product: "related to nucleosome remodeling	BX897674_24	1e-143	268.2	254.8	268.1	214.0	Myb-like DNA-binding domain, SWIRM domain	
	Pc22g15830	strong similarity to potential regulator of leucine permease gene expression Sac3 - Saccharomyces cerevisiae	probable leucine permease transcription regulator - fission yeast	T41415	4e-64	57.6	52.4	29.1	27.0	SAC3 GANP domain	
SART1	Pc22g13140	strong similarity to hypoxia-associated transcriptional activator SART1 - Tetraodon nigroviridis	hypothetical protein BQO2240 [imported] - Neurospora crassa	T49622	5e-92	150.6	148.9	158.0	126.7	SART1 family	
	Pc20g03020	weak similarity to hypoxia-associated transcriptional activator SART1 - Tetraodon nigroviridis	product: "probable transcriptional regulator, Arac family"	BX572600_256	2e-99	15.6	4.9	9.3	7.4	DJ1/PP2A family	
SEF1P	Pc20g00650	strong similarity to hnf-3f-like transcription factor sep1p - Schizosaccharomyces pombe	hnf-3f-like transcription factor homolog sep1 - fission yeast	J50500	8e-30	96.3	65.7	36.8	36.2	Forkhead domain	
	Pc12g03120	strong similarity to regulator protein Sin3 - Saccharomyces cerevisiae	gene: "B5K2.070"; product: "related to transcriptional activator Sin3p"; Aspergillus nidulans Sin3p gene, complete cds	BX42632_7	0.0	126.5	118.1	75.6	60.0	Paired amphipathic helix repeat (PAH), Histone deacetylase (HDAC)	
SIP3	Pc12g02400	weak similarity to Sin3 interacting protein Sin3 - Saccharomyces cerevisiae	product: "Sin3p"; Aspergillus nidulans Sin3p gene, complete cds	AY150563_1	0.0	169.1	141.4	160.7	181.2	PH domain	
	Pc22g04440	strong similarity to transcription regulator Sin7 - Saccharomyces cerevisiae	product: "Sin7p"; Aspergillus nidulans Sin7p gene, complete cds	AY168636_1	0.0	294.4	267.9	267.9	206.5	HSF-type DNA-binding domain, Response regulator receiver domain	
SKN7	Pc22g06090	weak similarity to transcription regulator Sin7 - Saccharomyces cerevisiae	Neurospora crassa DNA linkage group II BAC clone B18P7	BA25594_12	5e-05	75.6	68.8	67.0	55.4	No domain identified	
	Pc22g06380	weak similarity to transcription regulator Sin7 - Saccharomyces cerevisiae	gene: "OS.NB00365009.9"; product: "putative heat shock transcription factor"; Schizosaccharomyces pombe	AP033309_9	1e-05	38.5	36.4	22.9	11.0	HSF-type DNA-binding domain	
SNF2	Pc22g06380	strong similarity to SWI5N-related, matrix-associated, actin-dependent regulator of chromatin SMARCD1 - Homo sapiens	gene: "B234.1200"; product: "related to BSWF clone 60 KDa	NCB234_11	1e-109	236.2	243.9	179.3	184.7	SWIRM domain	
	Pc21g17380	strong similarity to transcription regulator Snf2 - Saccharomyces cerevisiae	probable transcription regulator - fission yeast (Schizosaccharomyces)	T37561	0.0	428.6	393.3	348.8	317.9	GLN domain, HSA domain, SNF2 family N-terminal domain, Helicase	
SNF5	Pc21g19220	transcription activator Snf5 - Saccharomyces cerevisiae	gene: "B1387.230"; product: "related to transcription factor smf2p";	BX08807_23	1e-108	188.5	161.1	173.3	138.3	SNF5-SMARCB1/Nr1	
	Pc12g11550	weak similarity to hypothetical transcription regulator SPAC11D3.07c - Schizosaccharomyces pombe	probable regulator protein YHR178w - yeast (Saccharomyces)	S48917	7e-06	18.9	38.9	44.8	60.4	Fungal specific transcription factor domain	
SPAC11D3.07c	Pc15g01850	weak similarity to hypothetical transcription regulator SPAC11D3.07c - Schizosaccharomyces pombe	probable transcription regulator, binuclear cluster zinc-finger - fission	T37518	3e-04	23.3	20.0	25.4	30.0	No domain identified	
	Pc16g02010	weak similarity to hypothetical transcription regulator SPAC11D3.07c - Schizosaccharomyces pombe	probable transcription regulator, binuclear cluster zinc-finger - fission	T37518	3e-04	23.3	20.0	25.4	30.0	No domain identified	
SPAC139.03	Pc16g12650	weak similarity to hypothetical transcription regulator SPAC11D3.07c - Schizosaccharomyces pombe	probable transcription regulator, binuclear cluster zinc-finger - fission	T37518	3e-04	23.3	20.0	25.4	30.0	No domain identified	
	Pc18g05390	similarity to hypothetical transcription regulator SPAC11D3.07c - Schizosaccharomyces pombe	probable transcription regulator, binuclear cluster zinc-finger - fission	T37518	3e-04	23.3	20.0	25.4	30.0	No domain identified	
SPAC139.03	Pc20g15350	weak similarity to hypothetical transcription regulator SPAC11D3.07c - Schizosaccharomyces pombe [putative sequencing error]	gene: "B11E5.150"; product: "related to transcriptional activator Mat3p";	BX42620_15	0.0	118.0	114.0	92.2	86.0	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
	Pc21g12210	weak similarity to hypothetical transcription regulator SPAC11D3.07c - Schizosaccharomyces pombe	gene: "B11E5.150"; product: "related to transcriptional activator Mat3p";	BX42620_15	3e-40	74.2	71.2	63.5	58.1	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
SPAC139.03	Pc21g16600	weak similarity to hypothetical transcription regulator SPAC11D3.07c - Schizosaccharomyces pombe	probable transcription regulator, binuclear cluster zinc-finger - fission	T37518	3e-04	23.3	20.0	25.4	30.0	No domain identified	
	Pc20g13810	similarity to hypothetical transcription regulator SPAC11D3.07c - Schizosaccharomyces pombe	gene: "B11E5.150"; product: "related to transcriptional activator Mat3p";	BX42620_15	1e-87	7.8	13.0	8.0	19.9	Fungal specific transcription factor domain	
SPAC139.03	Pc12g09550	similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	gene: "B5K2.250"; product: "conserved hypothetical protein";	BX42620_15	0.0	126.5	118.1	75.6	60.0	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
	Pc12g14990	weak similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	transcription activator protein acv-15 [imported] - Neurospora crassa	T46651	4e-40	39.2	37.1	44.0	45.3	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
SPAC139.03	Pc13g04660	similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	probable transcription activator - fission yeast (Schizosaccharomyces)	T37604	5e-22	24.7	48.8	13.9	15.4	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
	Pc13g11110	similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	gene: "B268.250"; product: "conserved hypothetical protein";	BX42620_15	2e-41	8.6	14.6	10.3	22.5	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
SPAC139.03	Pc13g12410	similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	probable transcription activator - fission yeast (Schizosaccharomyces)	T37604	1e-08	116.1	112.2	101.3	123.9	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
	Pc14g00280	weak similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	gene: "ZFR1"; product: "Zr1"; Gibberella moniliformis Zr1 (ZFR1)	AY493191_1	8e-08	12.3	10.6	8.3	12.5	Zn(II)(Cys) ₆	
SPAC139.03	Pc15g01860	similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	gene: "B1E8.250"; product: "conserved hypothetical protein";	BX40800_25	2e-82	17.2	18.4	21.9	20.3	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
	Pc16g00410	similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	probable transcription activator - fission yeast (Schizosaccharomyces)	T37604	5e-08	116.1	112.2	101.3	123.9	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
SPAC139.03	Pc16g00790	similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	gene: "B1E8.250"; product: "conserved hypothetical protein";	BX40800_25	2e-82	17.2	18.4	21.9	20.3	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
	Pc16g01070	similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	gene: "B1E8.250"; product: "conserved hypothetical protein";	BX40800_25	2e-82	17.2	18.4	21.9	20.3	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
SPAC139.03	Pc16g04810	similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	gene: "B1E8.250"; product: "conserved hypothetical protein";	BX40800_25	2e-82	17.2	18.4	21.9	20.3	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
	Pc16g07770	similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	gene: "B1E8.250"; product: "conserved hypothetical protein";	BX40800_25	2e-82	17.2	18.4	21.9	20.3	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
SPAC139.03	Pc16g08180	strong similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	gene: "B1E8.250"; product: "conserved hypothetical protein";	BX40800_25	2e-82	17.2	18.4	21.9	20.3	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
	Pc16g10100	similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	gene: "B1E8.250"; product: "conserved hypothetical protein";	BX40800_25	2e-82	17.2	18.4	21.9	20.3	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
SPAC139.03	Pc17g01190	weak similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	probable transcription activator - fission yeast (Schizosaccharomyces)	T37604	5e-07	122.5	123.3	13.0	1.8	Zn(II)(Cys) ₆ , Zn(II)(Cys) ₆	
	Pc18g03150	weak similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe [putative sequencing error]	probable transcription activator - fission yeast (Schizosaccharomyces)	T37604	1e-08	20.3	30.1	44.9	82.4	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
SPAC139.03	Pc18g03810	similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	probable transcription activator protein [imported] - fission yeast	T50198	3e-35	142.3	130.1	207.1	189.9	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
	Pc18g04010	similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	probable transcription activator - fission yeast (Schizosaccharomyces)	T37604	5e-08	116.1	112.2	101.3	123.9	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
SPAC139.03	Pc20g04260	weak similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	gene: "B11E23.130"; product: "conserved hypothetical protein";	NCB11E23_12	1e-48	40.1	33.8	37.9	43.1	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
	Pc20g11250	similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	probable transcription activator - fission yeast (Schizosaccharomyces)	T37604	2e-25	92.1	76.6	91.8	72.5	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
SPAC139.03	Pc20g12500	similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	gene: "B11E23.130"; product: "conserved hypothetical protein";	NCB11E23_12	1e-48	40.1	33.8	37.9	43.1	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
	Pc20g12500	similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	probable transcription activator - fission yeast (Schizosaccharomyces)	T37604	2e-25	92.1	76.6	91.8	72.5	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
SPAC139.03	Pc20g12500	similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	gene: "B11E23.130"; product: "conserved hypothetical protein";	NCB11E23_12	1e-48	40.1	33.8	37.9	43.1	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
	Pc20g12500	similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	probable transcription activator - fission yeast (Schizosaccharomyces)	T37604	2e-25	92.1	76.6	91.8	72.5	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
SPAC139.03	Pc20g12500	similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	gene: "B11E23.130"; product: "conserved hypothetical protein";	NCB11E23_12	1e-48	40.1	33.8	37.9	43.1	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
	Pc20g12500	similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	probable transcription activator - fission yeast (Schizosaccharomyces)	T37604	2e-25	92.1	76.6	91.8	72.5	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
SPAC139.03	Pc20g12500	similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	gene: "B11E23.130"; product: "conserved hypothetical protein";	NCB11E23_12	1e-48	40.1	33.8	37.9	43.1	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
	Pc20g12500	similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	probable transcription activator - fission yeast (Schizosaccharomyces)	T37604	2e-25	92.1	76.6	91.8	72.5	Zn(II)(Cys) ₆	Fungal specific transcription factor domain
SPAC139.03	Pc20g12500	similarity to hypothetical transcription activator SPAC139.03 - Schizosaccharomyces pombe	gene: "B11E23.130"; product: "conserved hypothetical protein";	NCB11E23_12	1e-48	40.1	33.8</				

SPT7	Pc13g09210	strong similarity to transcription factor Spt7 - Saccharomyces cerevisiae	gene: "B13D24.060": product: "related to transcription regulator SPT17"; regulatory protein SPT10 - yeast (Saccharomyces cerevisiae)	BX908789_6	0.0	7.2	6.8	18.9	9.7	Bromodomain, Bromo TP (bromodomain associated domain)
	Pc13g01610	strong similarity to regulator protein Spt10 - Saccharomyces cerevisiae	product: "suppressor protein spt23-related, with arginin repeats";	BX908789_6	7e-56	157.8	174.0	140.9	162.2	Acetyltransferase (GNAT) family, H2C2
SPT23	Pc13g13130	similarity to transcription suppressor protein Spt23 - Saccharomyces cerevisiae	product: "suppressor protein spt23-related, with arginin repeats";	BX908789_6	0.0	173.3	146.8	95.9	76.5	PT17C domain, Arginin repeat
	Pc13g03740	weak similarity to siderophore biosynthesis repressor SREA - Aspergillus nidulans	GATA-transcription factor - Penicillium chrysogenum	JC6170	8e-07	2.2	1.8	0.6	2.2	No domain identified
SREBP-1	Pc20g05890	similarity to sterol regulatory element binding protein 1 srebp-1 - Homo sapiens	gene: "B8624.060": product: "related to mvc-type bHLH transcription factor";	BX907676_6	4e-80	210.9	182.8	154.9	168.7	Helix-loop-helix DNA-binding domain
	Pc12g05640	GATA transcription factor snp1 - Penicillium chrysogenum	GATA-transcription factor - Penicillium chrysogenum	AF40308_4	3e-10	221.4	189.5	122.1	76.1	GATA zinc finger
STE11	Pc16g06680	similarity to transcription factor ste11p - Schizosaccharomyces pombe	gene: "mat1-2": product: "mating type 1-2 protein"; Mycoplasma pneumoniae	AF40308_4	3e-10	17.6	4.5	8.4	5.7	HMG (high mobility group) box
	Pc22g05580	similarity to transcription factor ste11p - Schizosaccharomyces pombe	product: "HMG-box protein STE11"; Pneumocystis carinii HMG-box	AY257481_1	7e-10	12.2	1.3	3.3	2.4	HMG (high mobility group) box
STU2	Pc12g07060	strong similarity to homeodomain DNA-binding transcription factor stu2 - Aspergillus nidulans	gene: "stu2": product: "homeodomain DNA-binding transcription factor";	AF080600_1	0.0	283.8	234.4	184.6	127.9	STE like transcription factor, C2H2
	Pc13g04920	strong similarity to transcription factor involved in differentiation stu2 - Aspergillus nidulans	gene: "SnA": product: "cell pattern formation-associated protein";	ENS1118_1	1e-143	255.5	192.3	125.0	69.0	APSES domain
SW4p	Pc13g08840	strong similarity to mating-type switching protein sw4p - Schizosaccharomyces pombe	gene: "MSH3": product: "msh3 homolog 3 (E. coli)"; Homo sapiens	AY275681_1	0.0	64.5	70.6	40.5	49.6	MuS domain I, II, III and V
	Pc13g09500	similarity to transcription factor Sw4b - Saccharomyces cerevisiae	hypothetical zinc-finger protein - fission yeast (Schizosaccharomyces pombe)	T30062	3e-22	305.3	251.1	166.3	80.0	C2H2
TAO3	Pc13g10900	strong similarity to transcription activator TAO3 - Saccharomyces cerevisiae	gene: "TAO3": product: "conserved hypothetical protein";	AF41242_1	0.0	142.0	101.5	108.6	142.8	No domain identified
	Pc13g12120	similarity to transcription activator thp1p - Schizosaccharomyces pombe	probable transcription regulator - fission yeast (Schizosaccharomyces pombe)	T40624	3e-08	4.1	1.6	4.0	5.5	Zn(II)/Cys6
TH1p	Pc13g04200	similarity to thiamine repressible gene th1p - Schizosaccharomyces pombe	gene: "tadB": product: "acetate regulatory DNA binding protein FadB";	ENL65097_1	4e-17	8.4	6.4	16.1	19.6	Zn(II)/Cys6, Fungal specific transcription factor domain
	Pc13g14910	weak similarity to thiamine repressible gene th1p - Schizosaccharomyces pombe	gene: "grr1": product: "zinc finger protein Grr1"; Schizosaccharomyces pombe	AF236387_1	1e-14	34.5	31.1	46.2	97.2	Zn(II)/Cys6, Fungal specific transcription factor domain
TH1p	Pc16g03680	similarity to thiamine repressible gene th1p - Schizosaccharomyces pombe	hypothetical protein SPB03.05 - fission yeast	T40621	8e-17	19.2	8.1	21.7	43.9	Zn(II)/Cys6, Fungal specific transcription factor domain
	Pc16g14530	similarity to thiamine-repressible protein th1p - Schizosaccharomyces pombe	gene: "rtr1+": product: "transcription factor"; Schizosaccharomyces pombe	SPMT1_X_1	3e-19	3.5	2.9	8.8	3.4	Zn(II)/Cys6, Fungal specific transcription factor domain
TH1p	Pc18g06190	similarity to thiamine repressible gene th1p - Schizosaccharomyces pombe	gene: "B11E5.150": product: "related to transcriptional activator Mut3p"; probable regulator protein - fission yeast (Schizosaccharomyces pombe)	BX846200_15	6e-89	38.5	31.8	41.5	41.9	Zn(II)/Cys6, Fungal specific transcription factor domain
	Pc20g02500	weak similarity to thiamine repressible gene th1p - Schizosaccharomyces pombe	probable regulator protein - fission yeast (Schizosaccharomyces pombe)	T36890	1e-15	36.5	33.3	23.4	44.2	Zn(II)/Cys6, Fungal specific transcription factor domain
TH1p	Pc12g12810	similarity to thiamine repressible gene th1p - Schizosaccharomyces pombe	gene: "B2E7.100": product: "related to lactose regulatory protein";	BX897675_10	1e-14	14.4	5.2	13.1	3.8	Zn(II)/Cys6, Fungal specific transcription factor domain
	Pc12g13820	similarity to thiamine-repressible protein th1p - Schizosaccharomyces pombe	gene: "B2E7.100": product: "related to lactose regulatory protein";	BX897675_10	1e-124	417.4	419.9	473.1	794.0	Zn(II)/Cys6, Fungal specific transcription factor domain
TH1p	Pc22g06450	similarity to thiamine-repressible protein th1p - Schizosaccharomyces pombe	thiamin repressible genes regulatory protein th1 [imported] - fission yeast	SA1962	1e-18	430.0	459.9	257.0	224.4	Zn(II)/Cys6, Fungal specific transcription factor domain
	Pc22g09250	similarity to thiamine-repressible protein th1p - Schizosaccharomyces pombe	gene: "rtr1+": product: "transcription factor"; Schizosaccharomyces pombe	SPMT1_X_1	6e-18	47.1	39.6	48.3	71.0	Zn(II)/Cys6, Fungal specific transcription factor domain
TH1p	Pc22g17630	similarity to thiamine repressible gene th1p - Schizosaccharomyces pombe	gene: "tadB": product: "tadB"; Aspergillus niger putative DNA binding	ANL56099_1	2e-24	9.9	10.3	9.3	13.0	Zn(II)/Cys6, Fungal specific transcription factor domain
TRI6	Pc22g05600	strong similarity to transcription activator Tri6 - Fusarium cerealis	gene: "TRI6": product: "regulatory protein"; Gibberella zeae strain	AY102589_3	8e-22	3.0	2.3	3.9	4.9	No domain identified
	Pc22g16130	similarity to transcription regulator TRI10 - Fusarium sporothricoides	gene: "TRI10": product: "regulatory protein"; Gibberella zeae strain	AY102584_5	1e-22	13.2	8.3	10.2	21.3	Zn(II)/Cys6
TRI6	Pc22g07140	strong similarity to positive regulator of purine utilisation uaf1 - Aspergillus nidulans	purine utilization positive regulator - Emmericella nidulans	SC4779	0.0	85.5	79.1	50.5	62.2	Zn(II)/Cys6, Fungal specific transcription factor domain
	Pc12g10080	weak similarity to positive regulator of purine utilisation uaf1 - Aspergillus nidulans	DNA centromeric region sequence from BAC DF26806, DP34F04,	CNS05045_32	3e-18	26.8	20.5	41.4	28.1	Zn(II)/Cys6
UAY	Pc13g04550	strong similarity to positive regulator of purine utilisation uaf1 - Aspergillus nidulans	probable membrane protein YLR278c - yeast (Saccharomyces cerevisiae)	SC0366	3e-38	29.3	25.9	24.7	28.8	Zn(II)/Cys6, Fungal specific transcription factor domain
	Pc15g01100	similarity to positive regulator of purine utilisation uaf1 - Aspergillus nidulans	gene: "PRF": product: "bacterial zinc transcription factor"; Nectria	AY218847_1	2e-26	12.0	1.7	14.6	29.0	Zn(II)/Cys6, Fungal specific transcription factor domain
UAY	Pc16g05770	weak similarity to positive regulator of purine utilisation uaf1 - Aspergillus nidulans [putative pseudogene]	zinc finger protein 1 homolog [imported] - Neurospora crassa	T49729	5e-09	218.1	161.9	119.7	80.8	Zn(II)/Cys6
	Pc16g13540	similarity to positive regulator of purine utilisation uaf1 - Aspergillus nidulans [putative sequencing error]	gene: "PRF": product: "bacterial zinc transcription factor"; Nectria	AY218847_1	2e-26	12.0	1.7	14.6	29.0	Zn(II)/Cys6, Fungal specific transcription factor domain
UAY	Pc20g03840	weak similarity to positive regulator of purine utilisation uaf1 - Aspergillus nidulans	purine utilization positive regulator - Emmericella nidulans	SC4779	4e-30	6.7	39.3	9.1	46.8	Zn(II)/Cys6, Fungal specific transcription factor domain
	Pc20g03700	strong similarity to positive regulator of purine utilisation uaf1 - Aspergillus nidulans	purine utilization positive regulator - Emmericella nidulans	SC4779	5e-34	3.6	0.7	2.0	24.0	Zn(II)/Cys6
UAY	Pc21g05090	weak similarity to positive regulator of purine utilisation uaf1 - Aspergillus nidulans	gene: "PRF": product: "bacterial zinc transcription factor"; Nectria	AY218847_1	2e-26	12.0	1.7	14.6	29.0	Zn(II)/Cys6, Fungal specific transcription factor domain
	Pc22g02710	weak similarity to positive regulator of purine utilisation uaf1 - Aspergillus nidulans	gene: "PRF": product: "bacterial zinc transcription factor"; Nectria	AY218847_1	2e-26	12.0	1.7	14.6	29.0	Zn(II)/Cys6, Fungal specific transcription factor domain
UGA3	Pc22g15880	weak similarity to regulator protein Uga3 - Saccharomyces cerevisiae	hypothetical fungal Zn(2)-Cys(6) zinc-finger protein - fission yeast	T41718	8e-12	283.3	209.8	160.3	134.5	Zn(II)/Cys6
	Pc18g04480	similarity to transcription factor Ume6 - Saccharomyces cerevisiae	gene: "rosa": product: "repressor of sexual development"; Aspergillus	ENR19682_1	1e-157	437.9	223.9	157.0	148.6	Zn(II)/Cys6
UM6	Pc16g01390	similarity to RNA polymerase II transcription factor Upc2 - Saccharomyces cerevisiae	gene: "B9B15.005": product: "putative protein"; Neurospora crassa	NCB9815_1	2e-11	0.6	2.2	1.8	2.4	Zn(II)/Cys6
	Pc12g04050	similarity to transcription activator Upc2 - Saccharomyces cerevisiae	probable membrane protein YDR213w - yeast (Saccharomyces cerevisiae)	SC1580	5e-08	10.7	10.7	12.7	12.7	Zn(II)/Cys6
UM6	Pc12g12360	weak similarity to hypothetical RNA polymerase II transcription factor Upc2 - Saccharomyces cerevisiae	gene: "B9B15.005": product: "putative protein"; Neurospora crassa	NCB9815_1	5e-16	7.0	4.7	10.2	7.2	Zn(II)/Cys6
	Pc13g01970	similarity to RNA polymerase II transcription factor Upc2 - Saccharomyces cerevisiae	gene: "pro1": product: "PRO1"; Neurospora crassa pro1 gene for	NCR238440_1	6e-06	180.8	178.7	96.6	63.2	Zn(II)/Cys6
UM6	Pc13g06330	similarity to transcription activator Upc2 - Saccharomyces cerevisiae	gene: "17E5.150": product: "conserved hypothetical protein";	NC17E5_14	2e-24	12.0	17.4	17.0	22.3	Zn(II)/Cys6
	Pc13g11250	similarity to transcription activator Upc2 - Saccharomyces cerevisiae	probable membrane protein YDR213w - yeast (Saccharomyces cerevisiae)	SC1580	4e-04	23.0	15.2	28.0	28.1	Zn(II)/Cys6
UM6	Pc14g00210	weak similarity to transcription activator Upc2 - Saccharomyces cerevisiae [putative sequencing error]	probable membrane protein YDR213w - yeast (Saccharomyces cerevisiae)	SC1580	3e-06	16.5	8.9	9.0	12.1	Zn(II)/Cys6
	Pc14g01000	similarity to transcription factor Upc2 - Saccharomyces cerevisiae	gene: "B9B15.005": product: "putative protein"; Neurospora crassa	NCB9815_1	1e-07	3.3	4.4	6.0	6.5	Zn(II)/Cys6
UM6	Pc14g01370	similarity to transcription factor Upc2 - Saccharomyces cerevisiae	probable membrane protein YDR213w - yeast (Saccharomyces cerevisiae)	SC1580	6e-08	94.7	144.9	235.0	330.1	Zn(II)/Cys6
	Pc16g02960	similarity to transcription activator Upc2 - Saccharomyces cerevisiae	gene: "B9B15.005": product: "putative protein"; Neurospora crassa	NCB9815_1	1e-18	29.9	29.7	39.3	53.5	Zn(II)/Cys6
UM6	Pc16g04030	weak similarity to RNA polymerase II transcription factor Upc2 - Saccharomyces cerevisiae	probable membrane protein YDR213w - yeast (Saccharomyces cerevisiae)	SC1580	8e-06	10.8	12.1	18.6	23.6	No domain identified
	Pc16g06330	weak similarity to transcription activator Upc2 - Saccharomyces cerevisiae [putative pseudogene]	zinc finger protein 1 homolog [imported] - Neurospora crassa	T49729	5e-09	218.1	161.9	119.7	80.8	Zn(II)/Cys6
UM6	Pc16g11730	similarity to transcription activator Upc2 - Saccharomyces cerevisiae	gene: "B9B15.005": product: "putative protein"; Neurospora crassa	NCB9815_1	2e-14	11.4	15.1	16.5	18.6	Zn(II)/Cys6
	Pc18g06880	weak similarity to RNA polymerase II transcription factor Upc2 - Saccharomyces cerevisiae	probable membrane protein YDR213w - yeast (Saccharomyces cerevisiae)	SC1580	1e-04	4.3	10.1	2.9	7.5	Zn(II)/Cys6
UM6	Pc20g05430	weak similarity to hypothetical RNA polymerase II transcription factor Upc2 - Saccharomyces cerevisiae	gene: "B9B15.005": product: "putative protein"; Neurospora crassa	NCB9815_1	6e-13	78.8	88.1	98.1	99.0	Zn(II)/Cys6
	Pc21g08710	weak similarity to transcription activator Upc2 - Saccharomyces cerevisiae	probable membrane protein YDR213w - yeast (Saccharomyces cerevisiae)	SC1580	2e-12	80.0	65.3	68.8	33.9	Zn(II)/Cys6
UM6	Pc21g09290	similarity to transcription activator Upc2 - Saccharomyces cerevisiae	zinc finger protein 1 homolog [imported] - Neurospora crassa	T49729	5e-12	74.0	40.1	67.8	52.6	Zn(II)/Cys6
	Pc21g12280	similarity to RNA polymerase II transcription factor Upc2 - Saccharomyces cerevisiae	probable membrane protein YDR213w - yeast (Saccharomyces cerevisiae)	SC1580	2e-12	80.0	65.3	68.8	33.9	Zn(II)/Cys6
UM6	Pc21g15030	similarity to transcription activator Upc2 - Saccharomyces cerevisiae	gene: "PRO1": product: "PRO1 protein"; Sordaria macrospora pro1	SAJ1051_1	6e-06	13.1	13.9	38.2	50.1	Zn(II)/Cys6
	Pc21g17780	weak similarity to transcription activator Upc2 - Saccharomyces cerevisiae	gene: "17E5.150": product: "conserved hypothetical protein";	NC17E5_14	1e-16	78.6	70.3	82.0	72.5	Zn(II)/Cys6
UM6	Pc21g18560	similarity to RNA polymerase II transcription factor Upc2 - Saccharomyces cerevisiae	hypothetical fungal Zn(2)-Cys(6) zinc-finger protein - fission yeast	T41718	2e-08	54.0	44.8	38.8	33.7	Zn(II)/Cys6
	Pc21g1670	weak similarity to hypothetical RNA polymerase II transcription factor Upc2 - Saccharomyces cerevisiae	probable membrane protein YLR228c - yeast (Saccharomyces cerevisiae)	SC1451	2e-07	2.8	1.0	3.9	5.1	Zn(II)/Cys6
UM6	Pc22g00480	similarity to transcription activator Upc2 - Saccharomyces cerevisiae	gene: "B9B15.005": product: "putative protein"; Neurospora crassa	NCB9815_1	4e-15	38.3	40.5	84.6	87.3	Zn(II)/Cys6
	Pc22g06200	similarity to transcription activator Upc2 - Saccharomyces cerevisiae	gene: "B9B15.005": product: "putative protein"; Neurospora crassa	NCB9815_1	6e-11	24.8	21.3	30.2	38.9	Zn(II)/Cys6
UM6	Pc22g07590	similarity to transcription activator Upc2 - Saccharomyces cerevisiae	probable membrane protein YDR213w - yeast (Saccharomyces cerevisiae)	SC1580	2e-10	30.1	29.7	36.6	40.8	No domain identified
	Pc22g16910	similarity to transcription activator Upc2 - Saccharomyces cerevisiae	product: "hypothetical protein"; Aspergillus fumigatus BAC pilot project	BX649605_122	2e-97	87.7	112.8	128.8	130.1	Zn(II)/Cys6
UM6	Pc22g20530	weak similarity to hypothetical RNA polymerase II transcription factor Upc2 - Saccharomyces cerevisiae	probable membrane protein YLR228c - yeast (Saccharomyces cerevisiae)	SC1451	4e-05	18.2	20.2	41.4	75.6	No domain identified
	Pc22g22500	weak similarity to hypothetical RNA polymerase II transcription factor Upc2 - Saccharomyces cerevisiae	probable membrane protein YLR228c - yeast (Saccharomyces cerevisiae)	SC1451	7e-06	52.0	35.9	82.8	72.3	No domain identified
UM6	Pc22g26430	similarity to transcription activator Upc2 - Saccharomyces cerevisiae	gene: "CAT8": product: "Cat8"; Candida albicans Ngt1 (NTG1) gene,	AF222088_3	3e-07	5.6	6.7	9.0	10.5	Zn(II)/Cys6
	Pc24g00540	weak similarity to transcription activator Upc2 - Saccharomyces cerevisiae	probable membrane protein YDR213w - yeast (Saccharomyces cerevisiae)	SC1580	8e-06	15.2	10.2	21.5	28.7	No domain identified
USF2	Pc22g15870	weak similarity to upstream stimulatory factor of transcription USF2 - Homo sapiens	gene: "penR2": product: "PENR2 protein"; Emmericella nidulans penR2	ENR17331_1	2e-50	406.4	337.3	221.6	212.6	Helix-loop-helix DNA-binding domain
	Pc16g03680	strong similarity to central regulator of blue light responses wcc1 - Neurospora crassa	gene: "traA": product: "GATA factor"; Emmericella nidulans GATA factor	AF316628_1	1e-172	32.8	33.0	29.5	71.9	PAS 3 domain, GATA domain
WETA	Pc22g03220	regulator protein wetaA - Penicillium chrysogenum	gene: "wetaA": product: "GATA factor"; Penicillium chrysogenum	SA6680	0.0	10.3	4.2	14.9	13.6	No domain identified
	Pc22g00710	similarity to transcription repressor Xbp1 - Saccharomyces cerevisiae	probable membrane protein YL101c - yeast (Saccharomyces cerevisiae)	SA4871	7e-05	9.9	6.1	12.8	11.3	No domain identified
XBP1	Pc21g02790	strong similarity to xylanolytic transcriptional activator xbrR - Aspergillus niger	gene: "xbrR": product: "transcription activator"; Aspergillus niger	AB042843_1	0.0	159.2	158.8	118.8	150.3	Zn(II)/Cys6, Fungal specific transcription factor domain
	Pc22g02960	strong similarity to xylanolytic transcriptional activator xbrR - Aspergillus niger	gene: "xbrR": product: "transcription activator"; Aspergillus niger	ANL1999_1	1e-136	118.4	93.8	90.5	81.6	Zn(II)/Cys6, Fungal specific transcription factor domain
YAP1	Pc06g00310	weak similarity to transcription factor YAP1 - Kluyveromyces fragilis	gene: "xbrR": product: "transcription activator"; Aspergillus niger	AB042843_1	0.0	159.2	1			

**Supplementary Table 27. Probe set characteristics of
DSM_PENa520255F, hybridized with Wis54-1255 genomic
DNA**

	#	%	Average Signal
Total Probe Sets:	15525		107.7
Number Present:	15346	98.80%	108.9
Number Absent:	158	1.00%	3
Number Marginal:	21	0.10%	14.5
 Annotated ORFs probe sets	 13704		 105
Number Present:	13615	99.40%	-
Number Absent:	76	0.60%	-
Number Marginal:	13	0.10%	-